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(54) **FLAGELLIN RELATED POLYPEPTIDES AND USES THEREOF**(71) Applicant: **Cleveland Clinic Foundation**, Cleveland, OH (US)(72) Inventors: **Andrei V. Gudkov**, East Aurora, NY (US); **Joseph A. DiDonato**, Westlake, OH (US); **Vadim Krivokrysenko**, Orchard Park, NY (US)(73) Assignee: **Cleveland Clinic Foundation**, Cleveland, OH (US)

(*) Notice: Subject to any disclaimer, the term of this patent is extended or adjusted under 35 U.S.C. 154(b) by 0 days.

(21) Appl. No.: **14/828,111**(22) Filed: **Aug. 17, 2015****Related U.S. Application Data**

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(51) **Int. Cl.**

A61K 38/16	(2006.01)
A61K 39/112	(2006.01)
C07K 14/255	(2006.01)
A61K 45/06	(2006.01)

(52) **U.S. Cl.**

CPC	C07K 14/255 (2013.01); A61K 38/164 (2013.01); A61K 39/0275 (2013.01); A61K 45/06 (2013.01)
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(58) **Field of Classification Search**

None

See application file for complete search history.

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Primary Examiner — S. Devi(74) *Attorney, Agent, or Firm — Morgan, Lewis & Bockius LLP*(57) **ABSTRACT**

The use of flagellin and flagellin related polypeptides for the protection of mammals from the effects of apoptosis is described.

8 Claims, 105 Drawing Sheets

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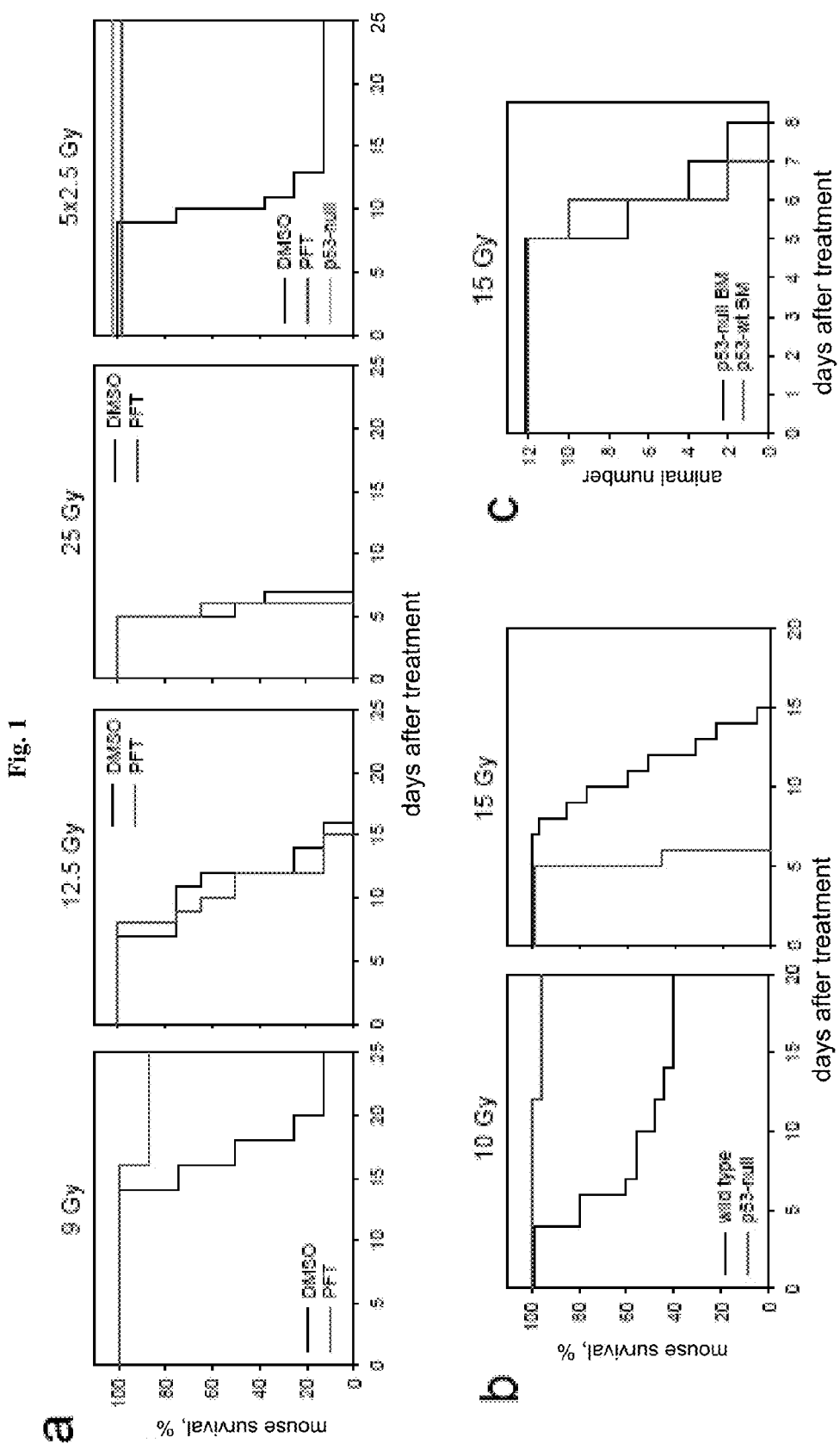


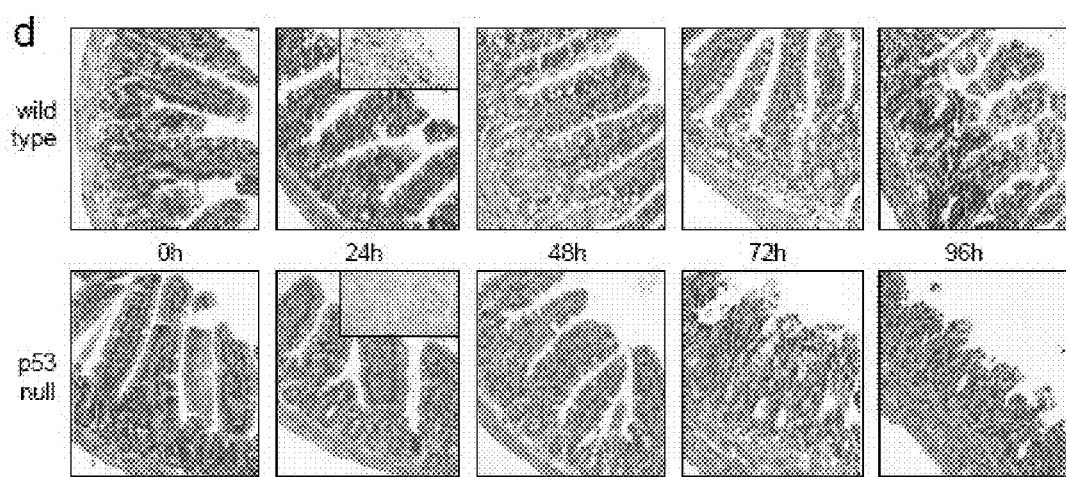
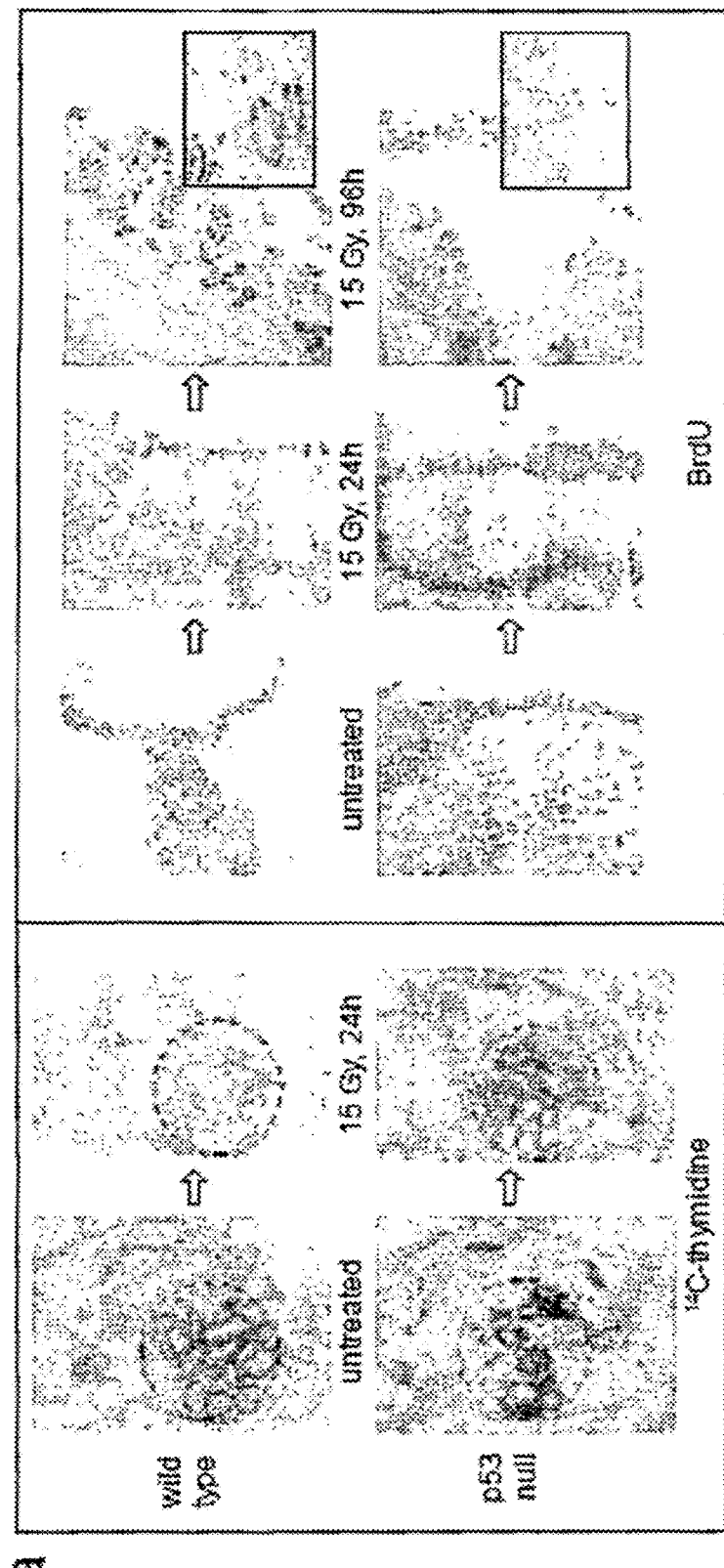
Fig. 1, CONTINUED

Fig. 2



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Fig. 2, continued

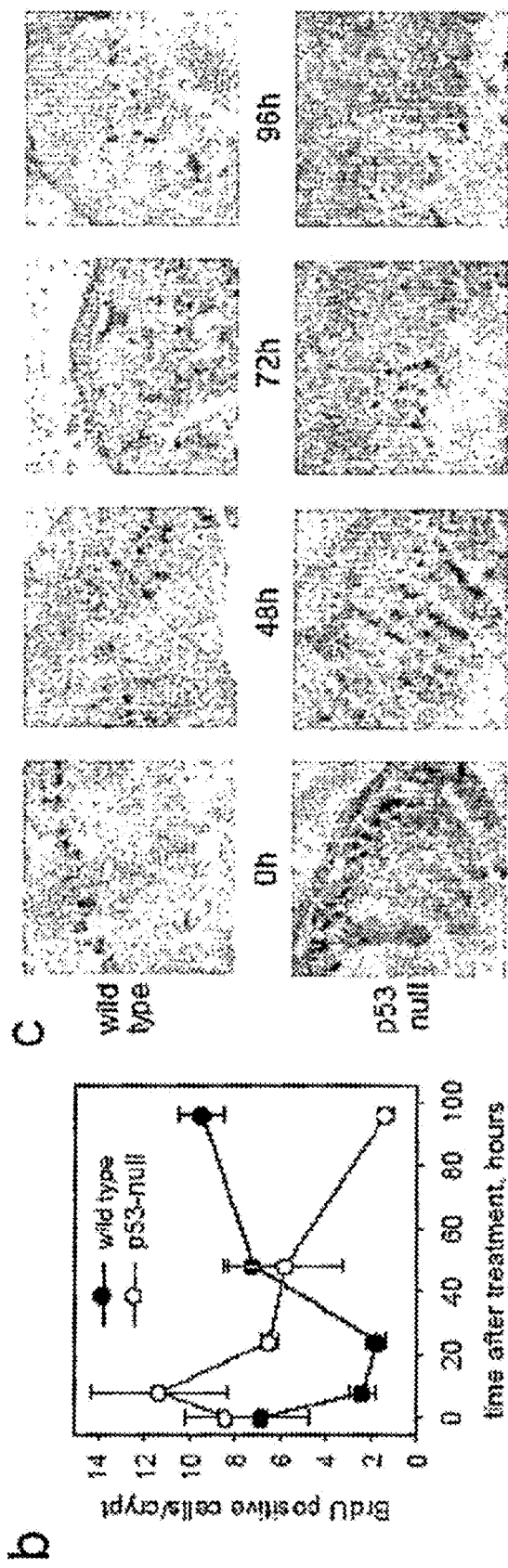


Fig. 3

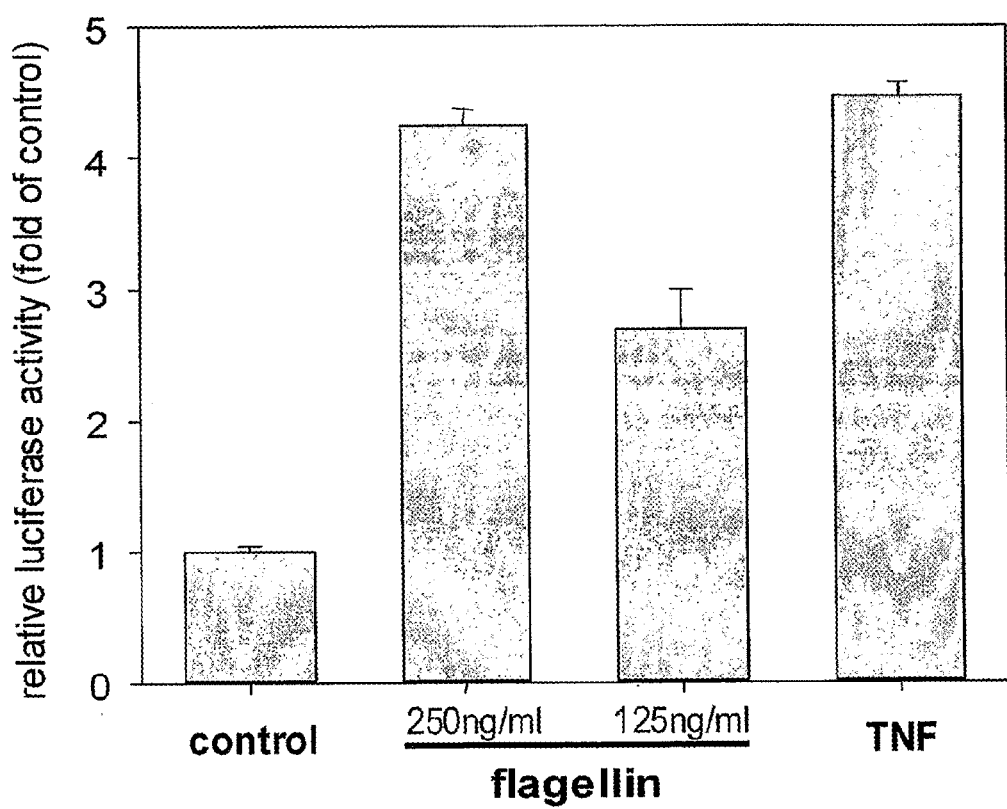


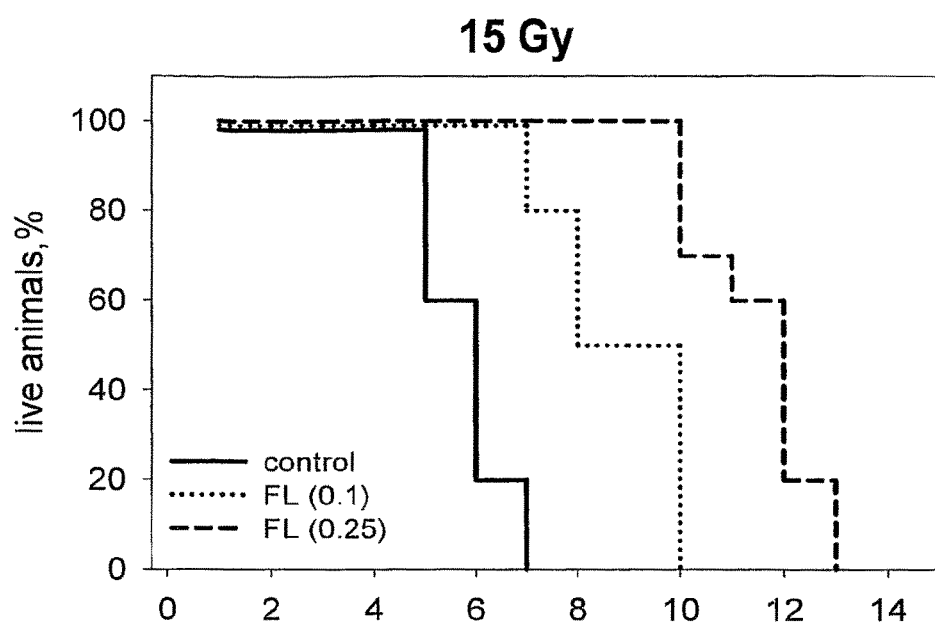
Fig. 4

Fig. 5

Small intestine, day 7

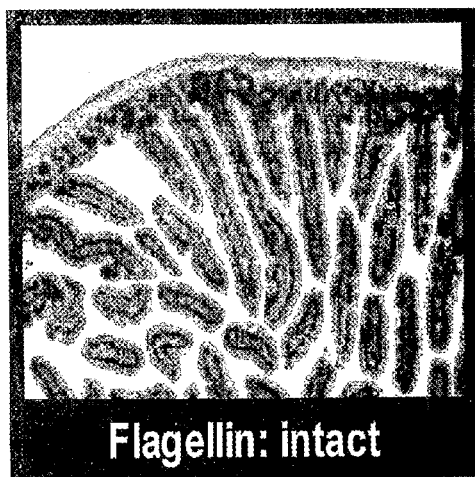
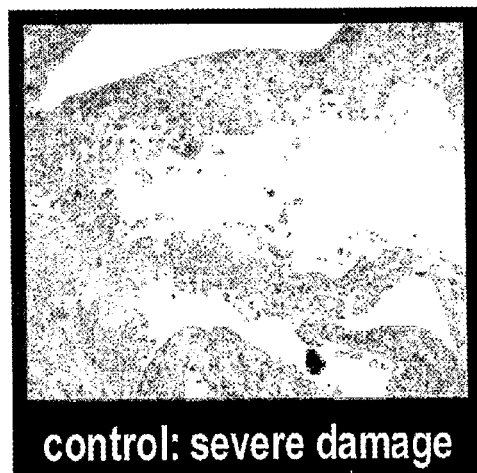


Fig. 6

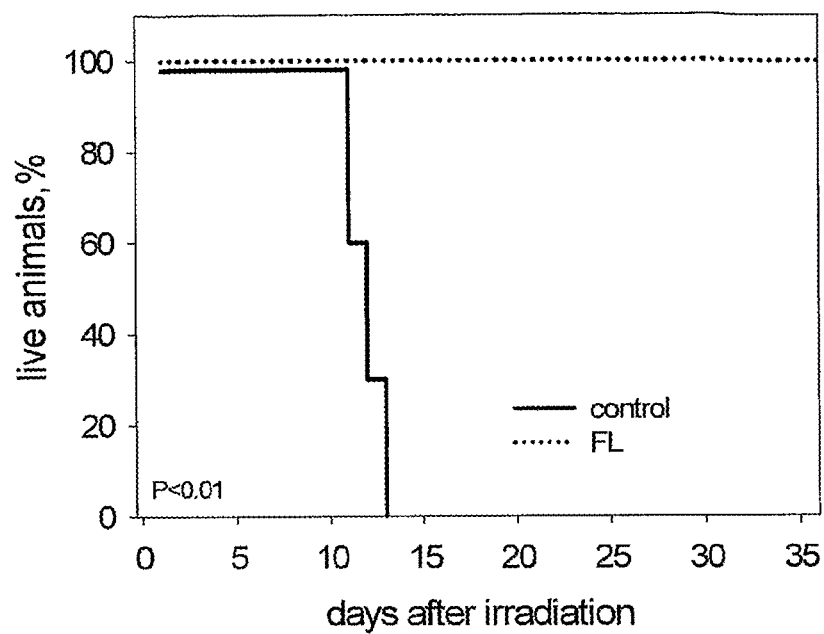
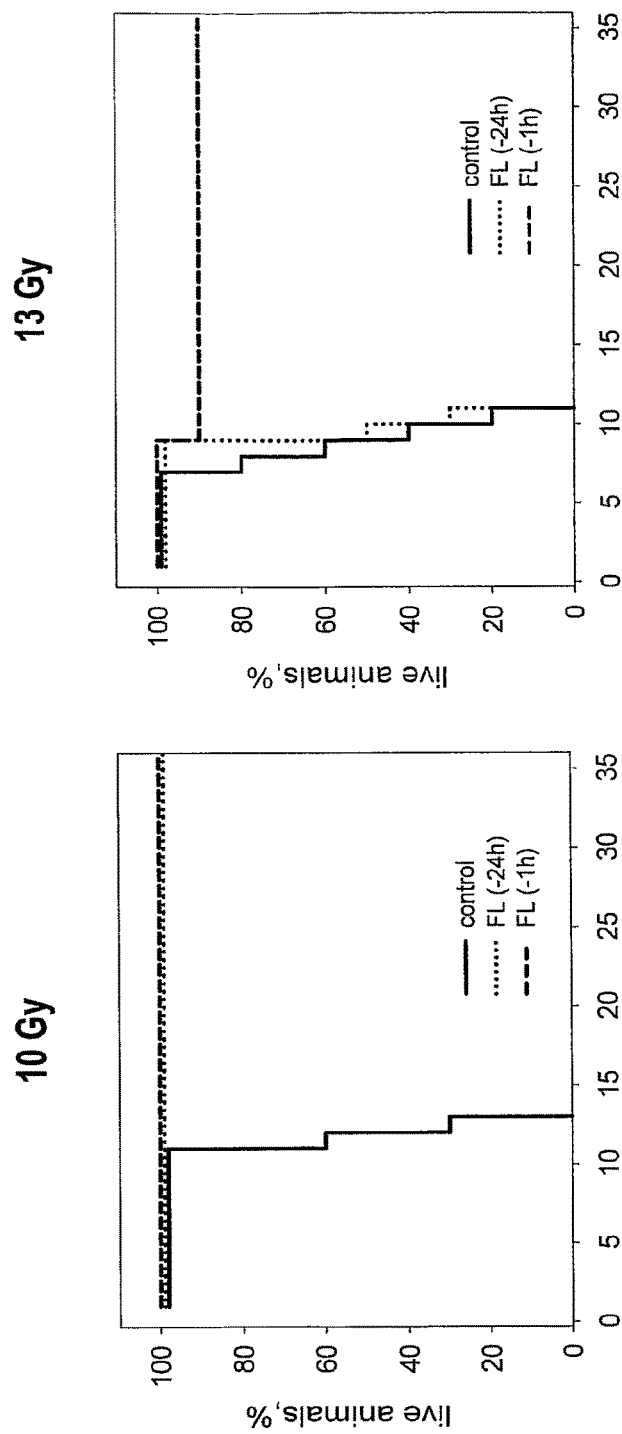


Fig. 7



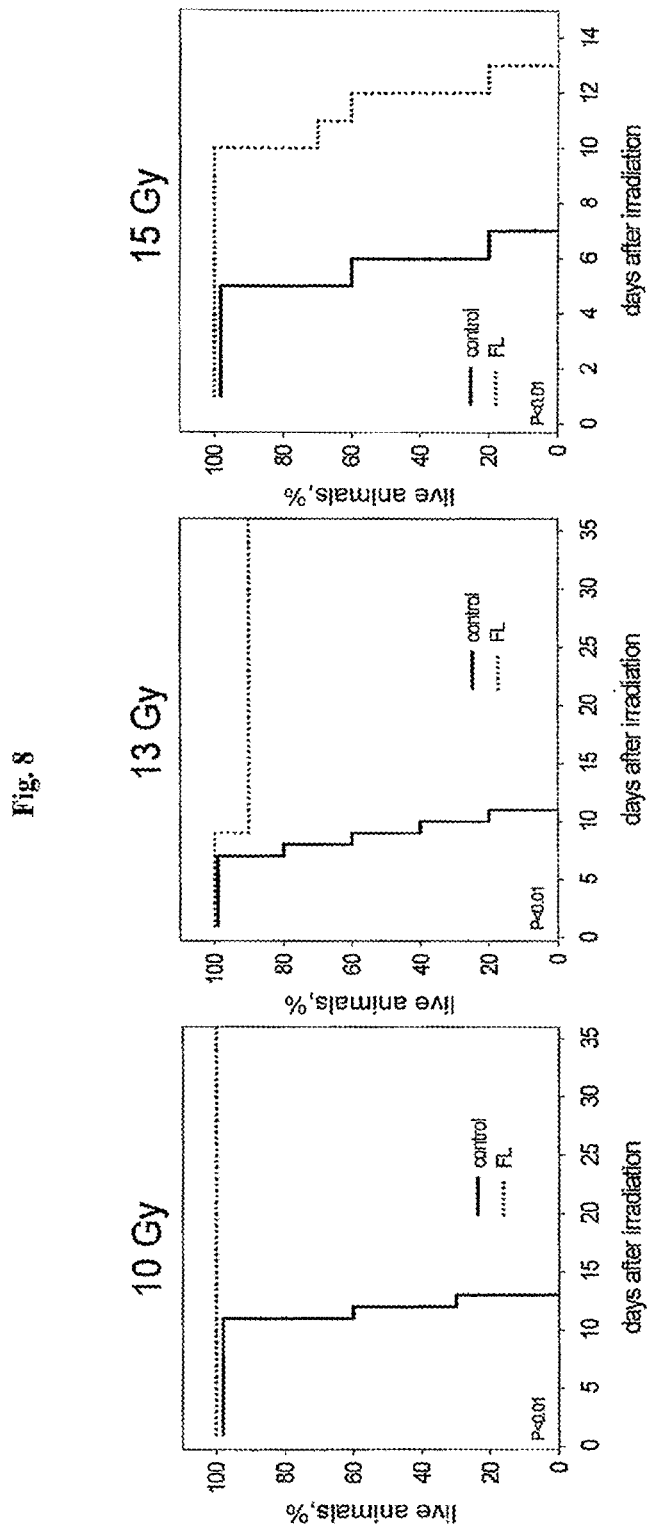


Fig. 9

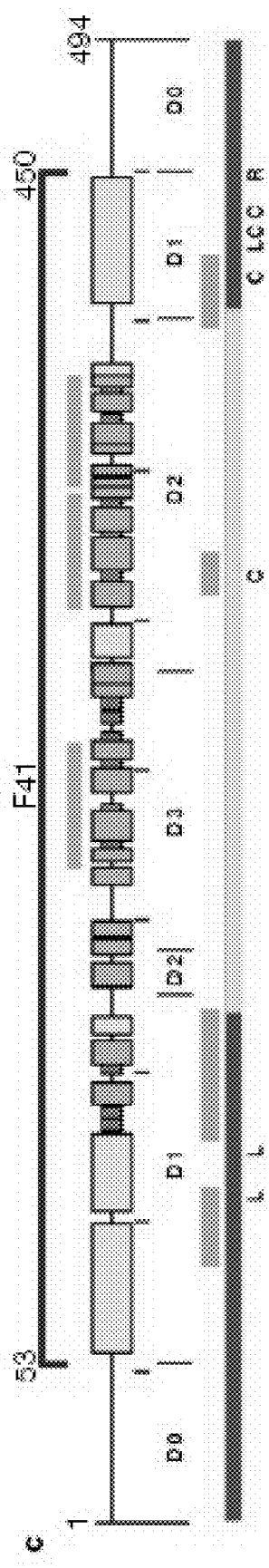


Fig. 10

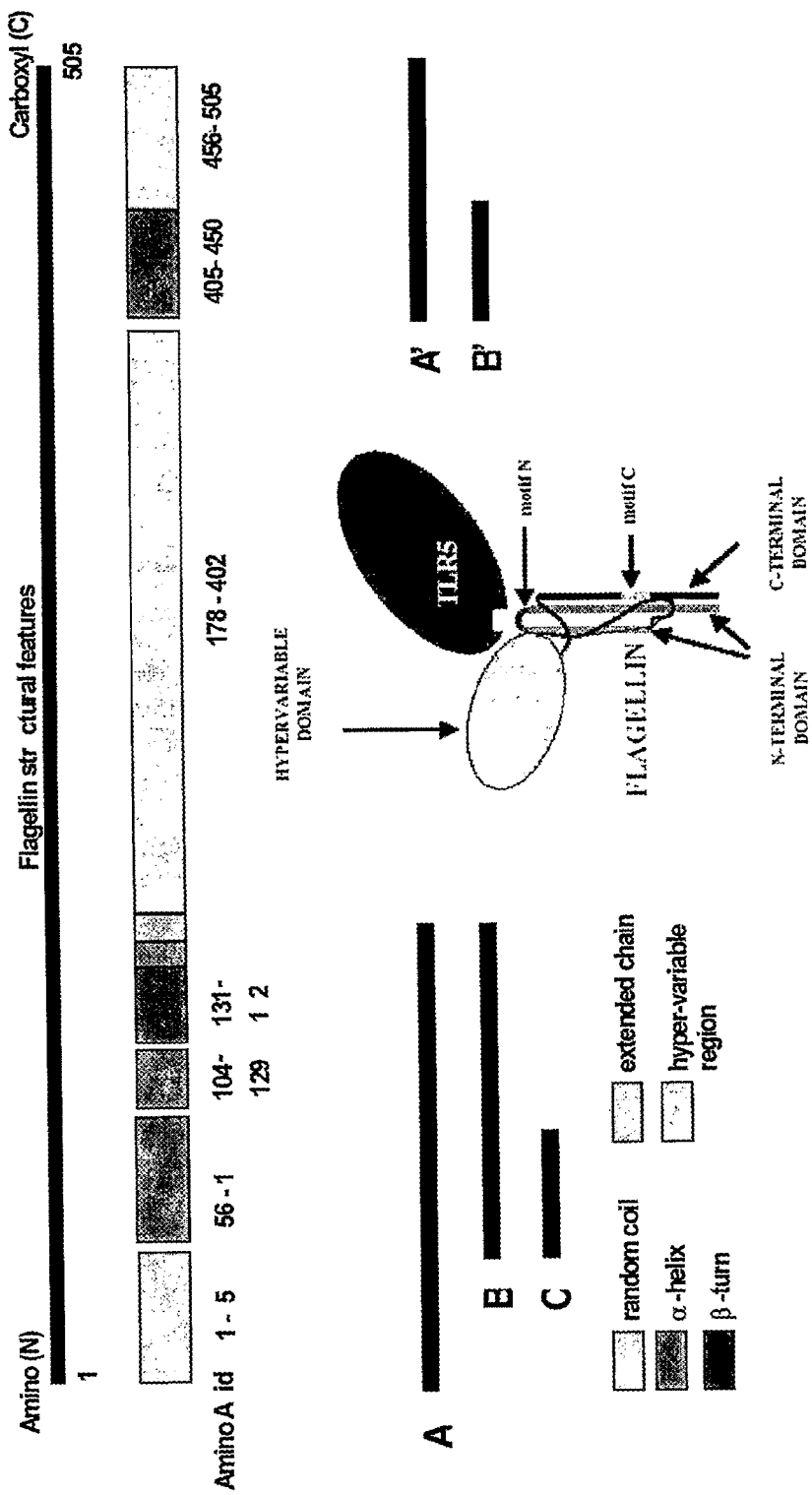


Fig. 11

MW HiC AA' BB'



Fig. 12

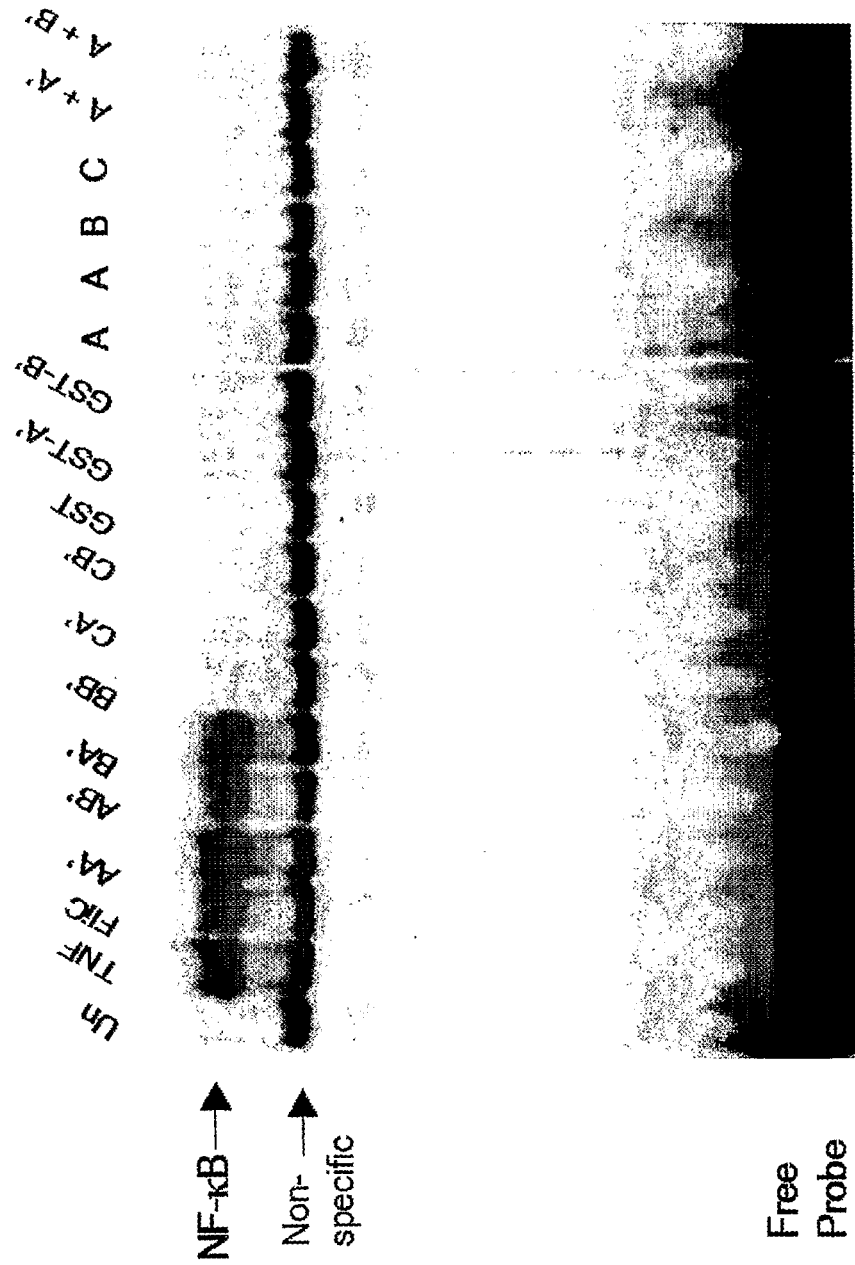


Fig. 13

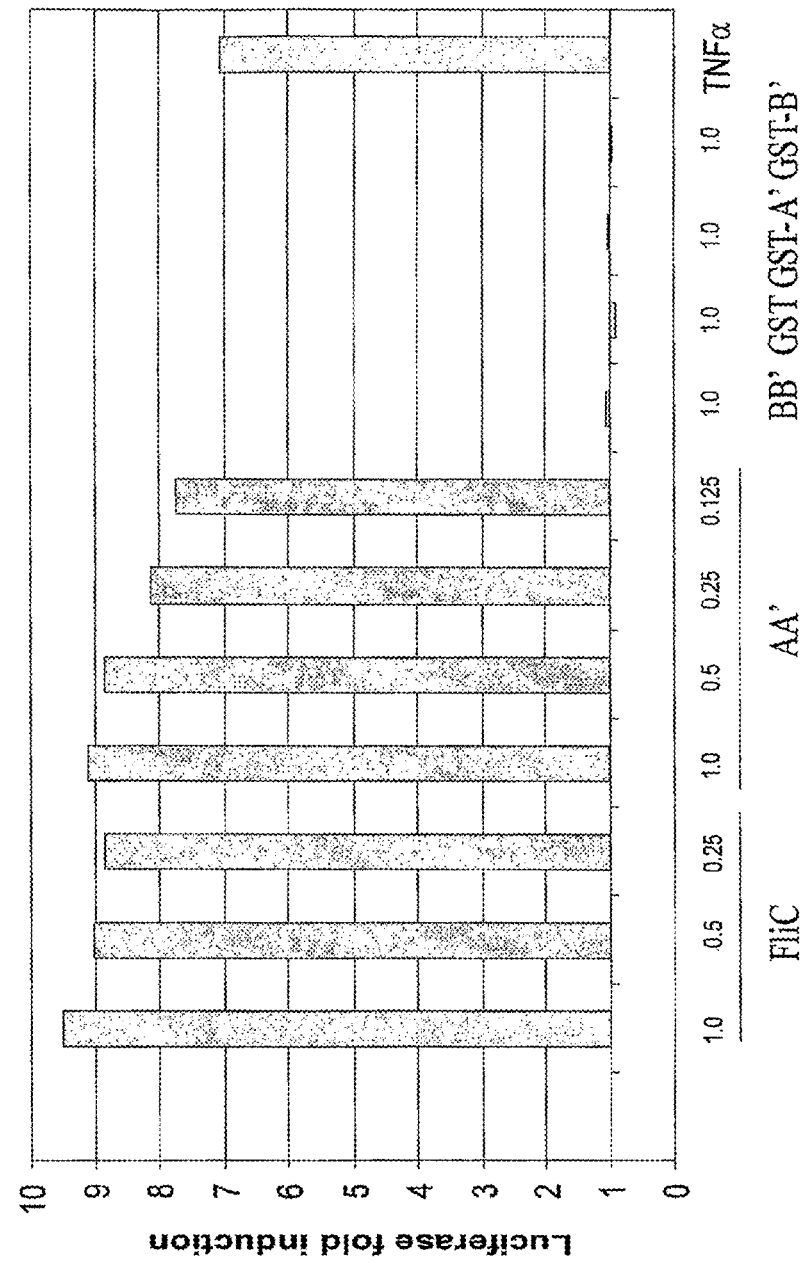


Fig. 14

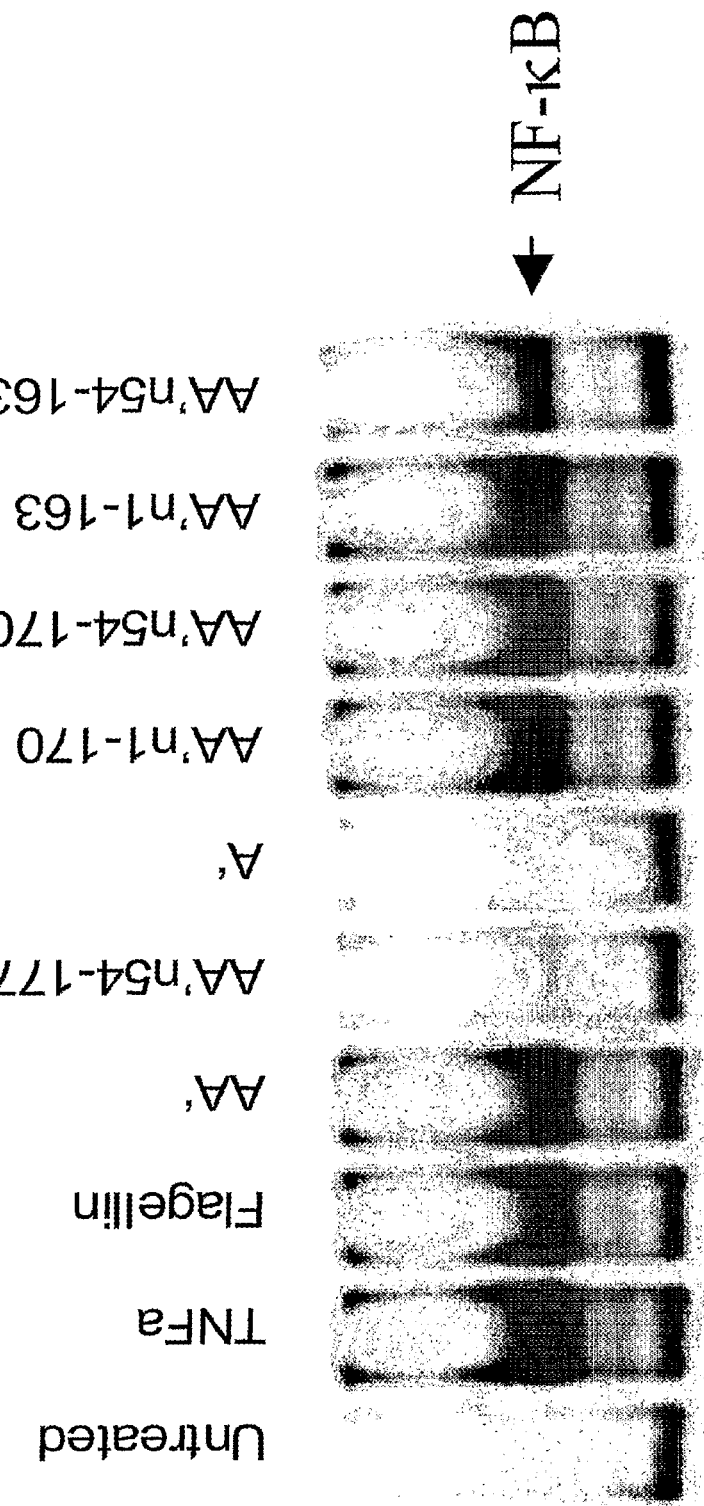
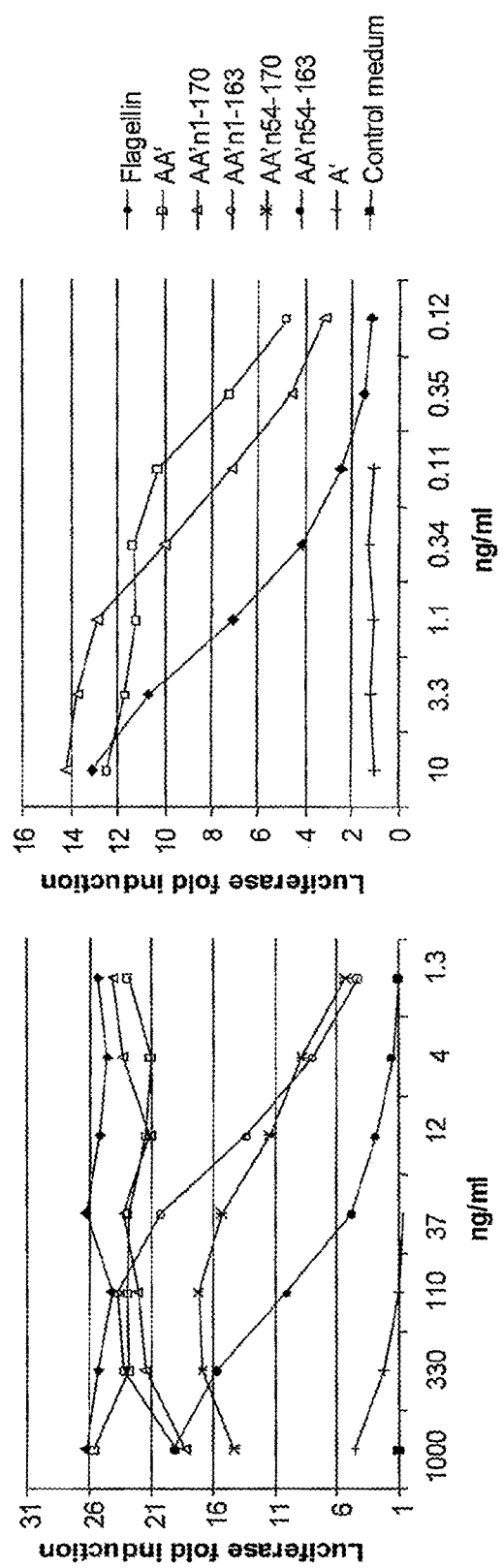


Fig. 15



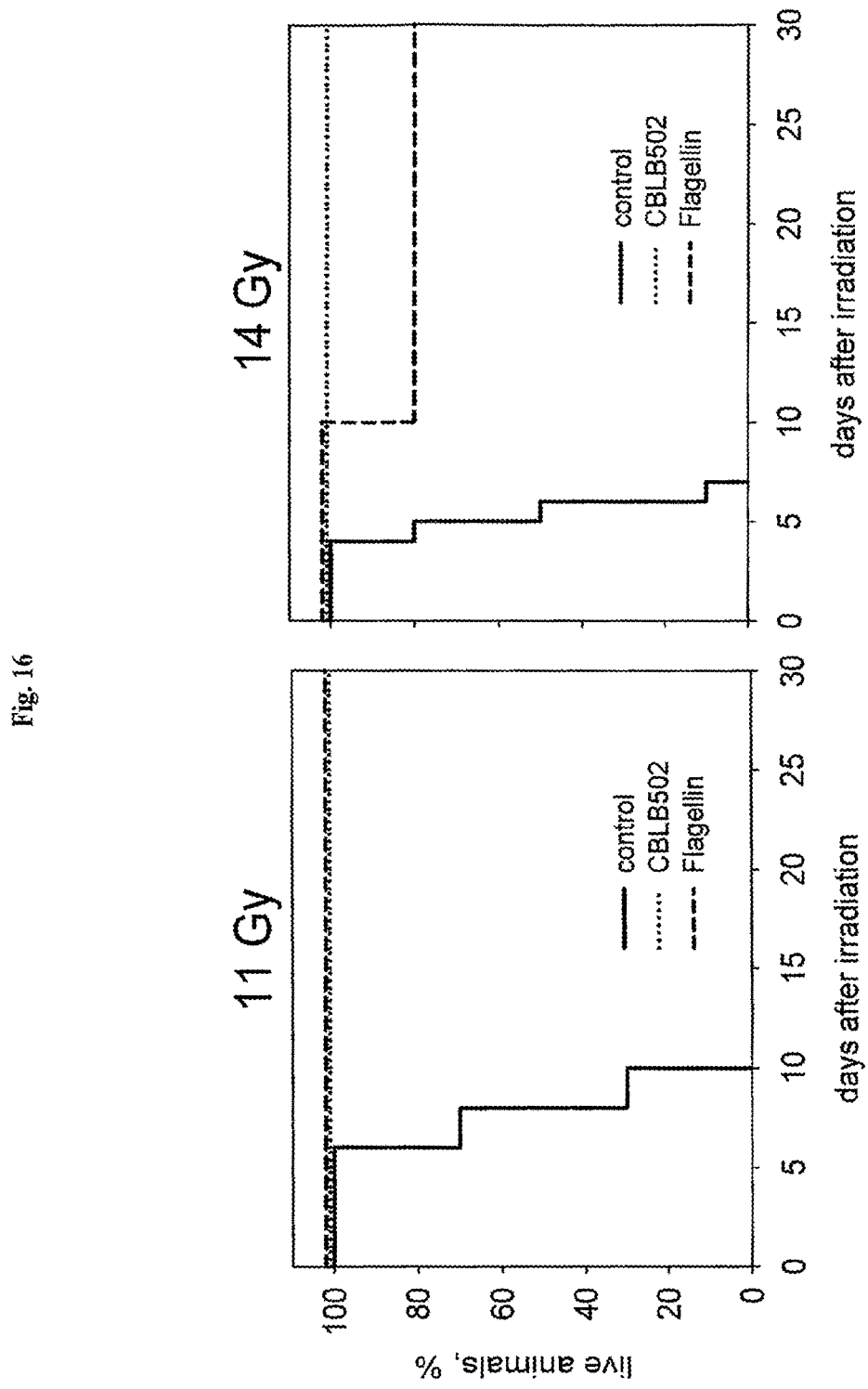


Fig. 17

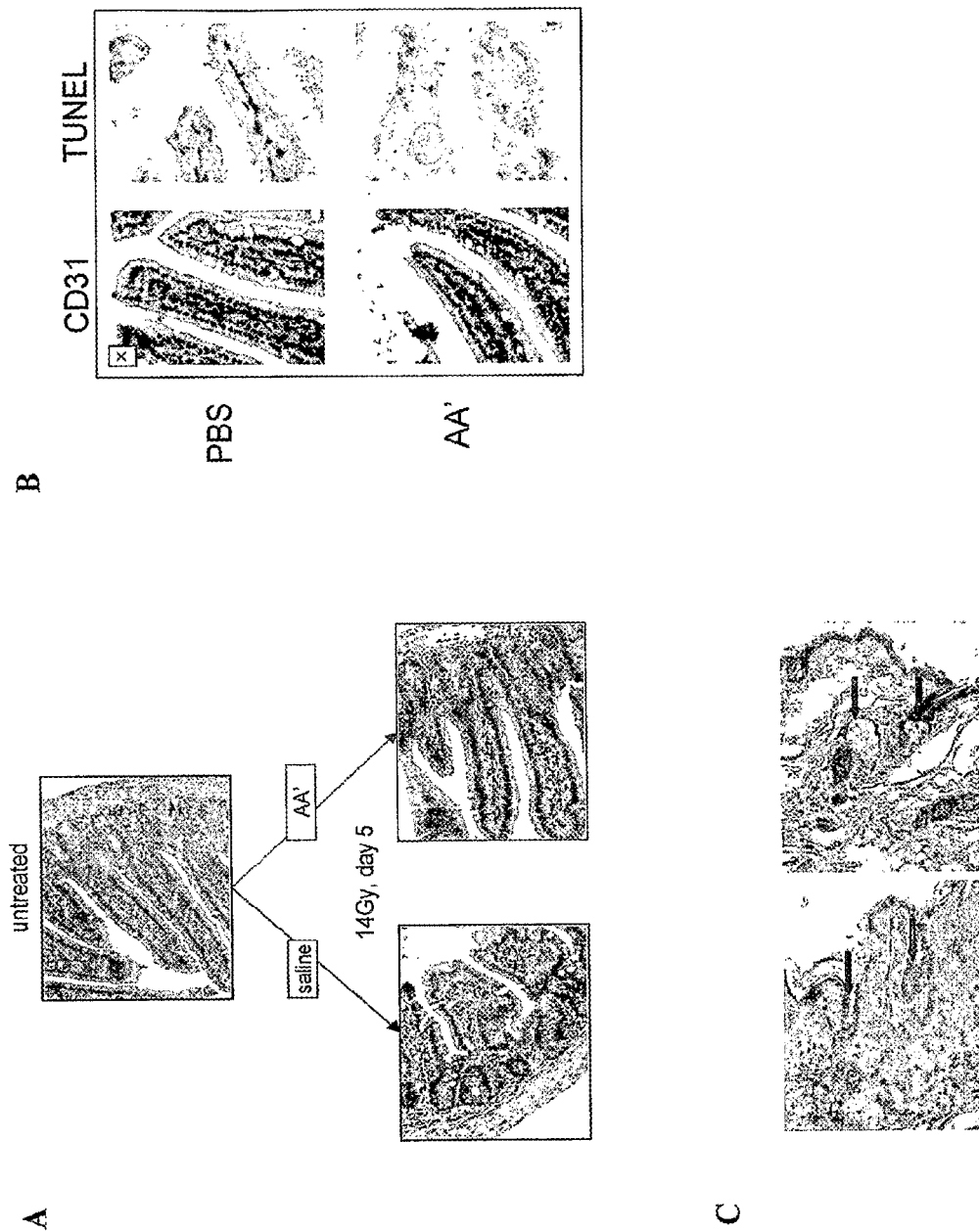


Fig. 18

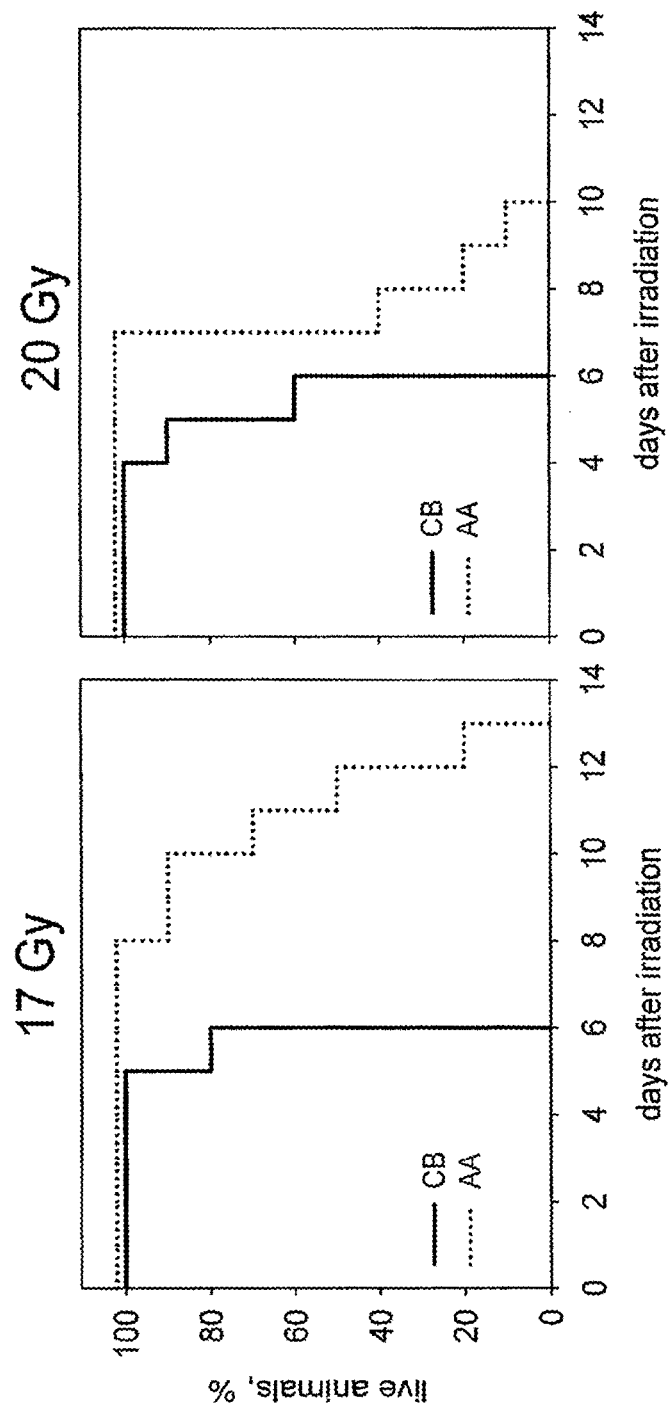


Fig. 19

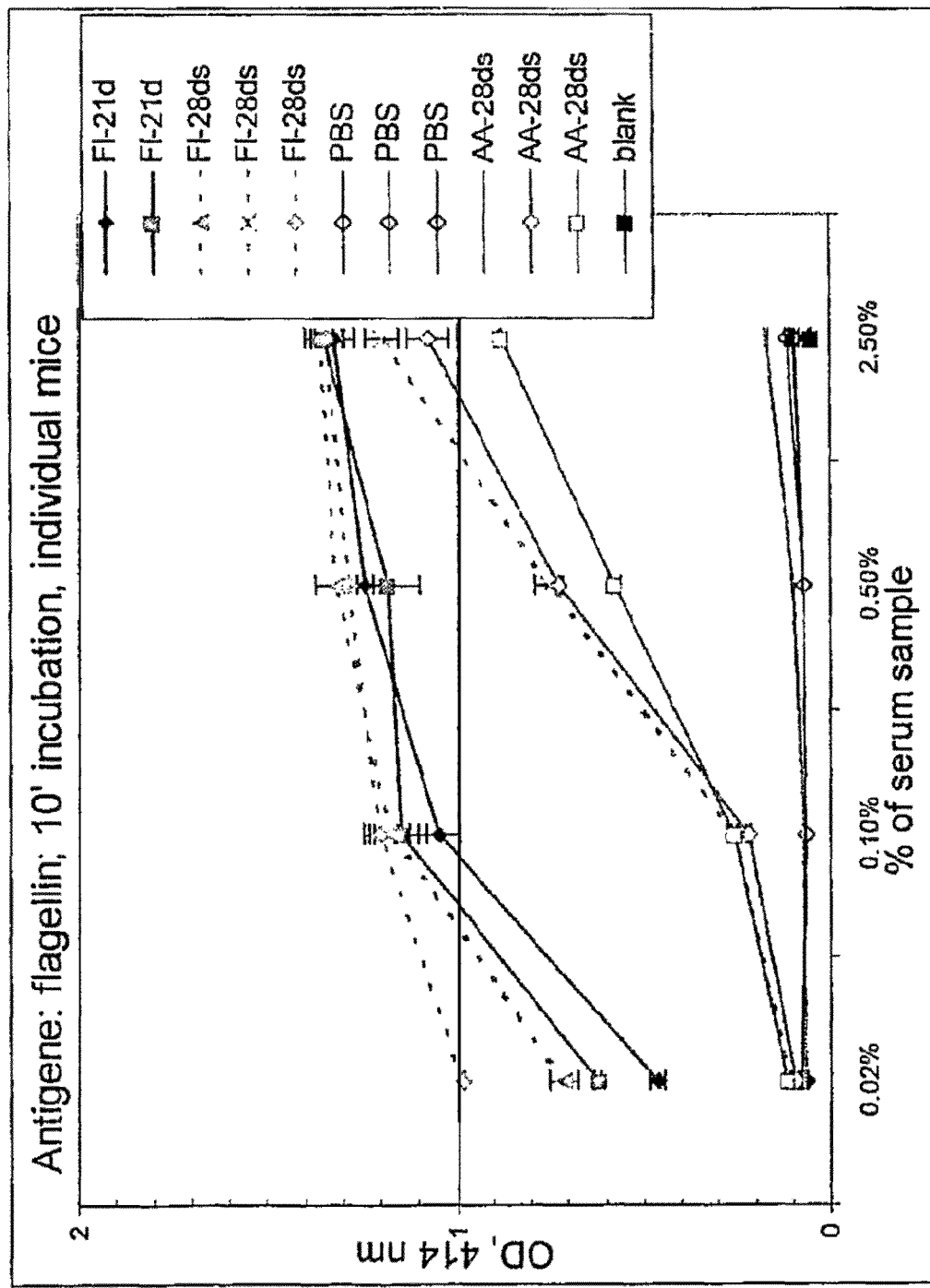


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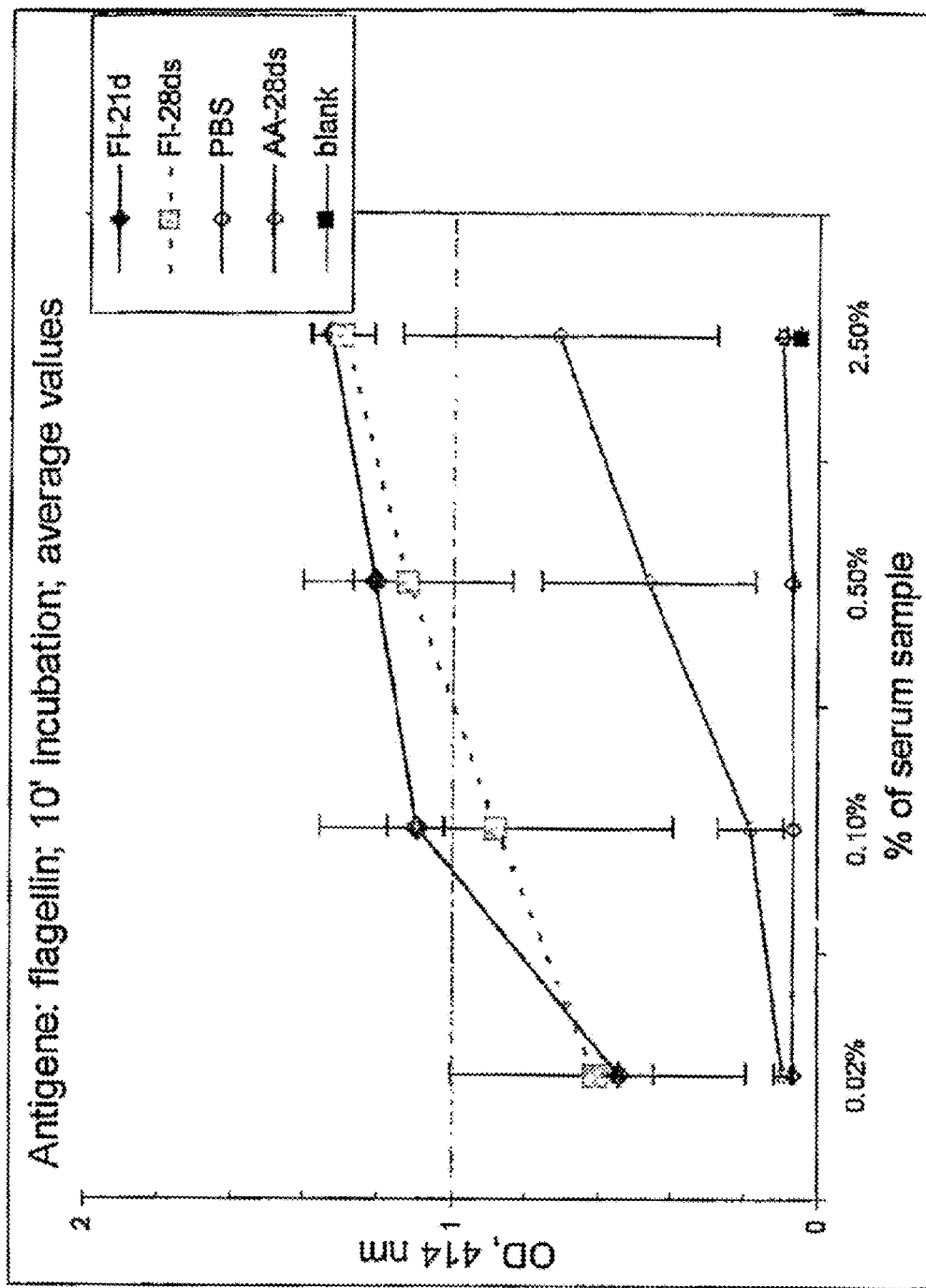


Fig. 20

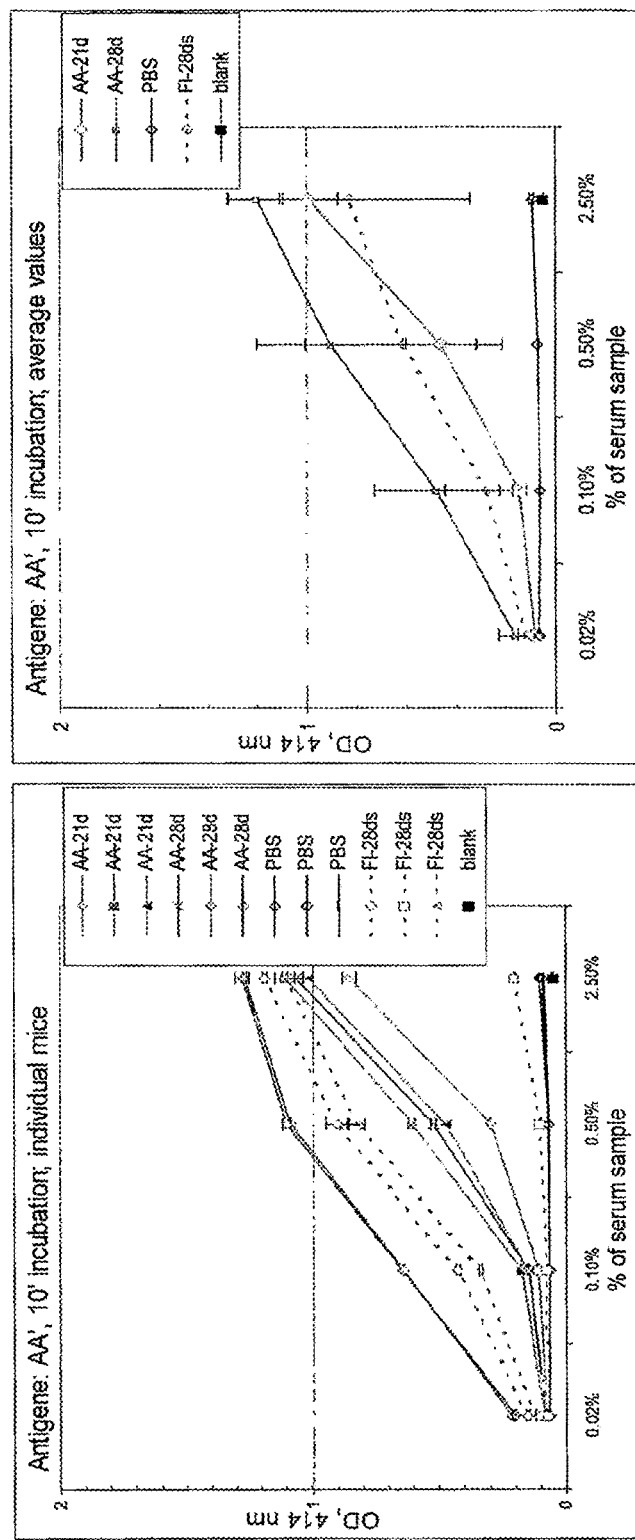


Fig. 21

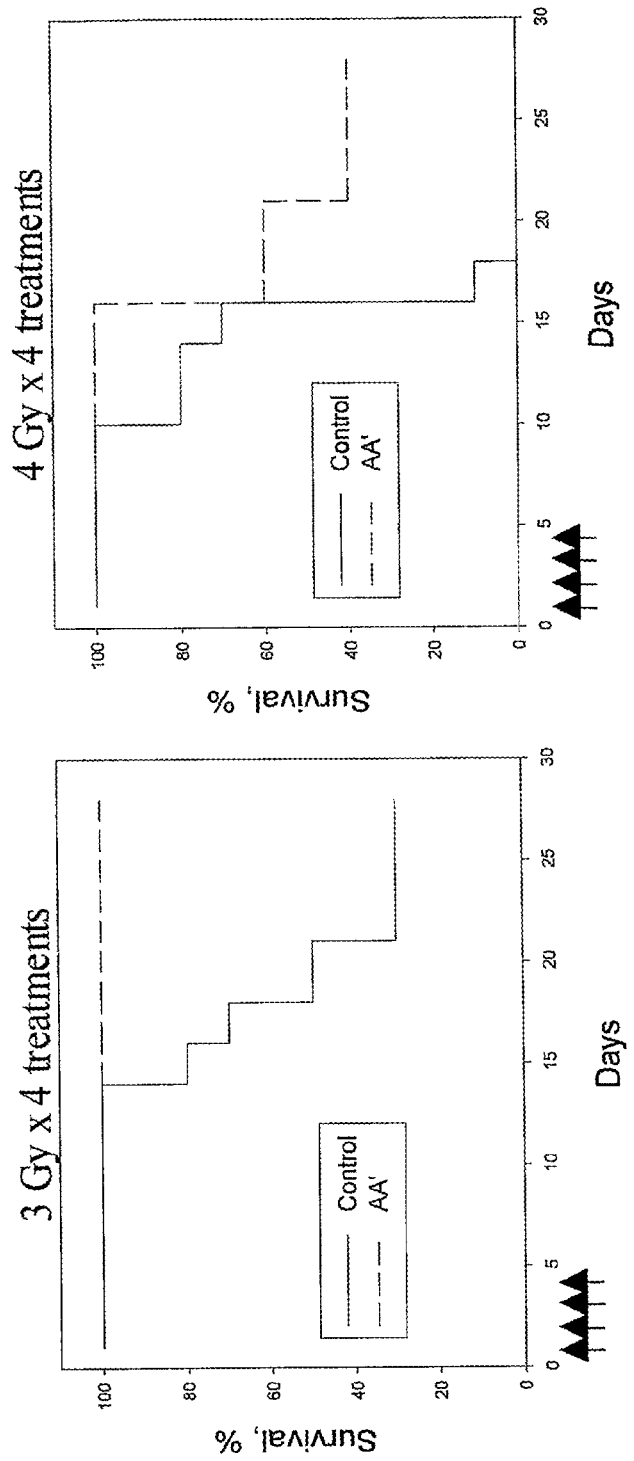


Fig. 22

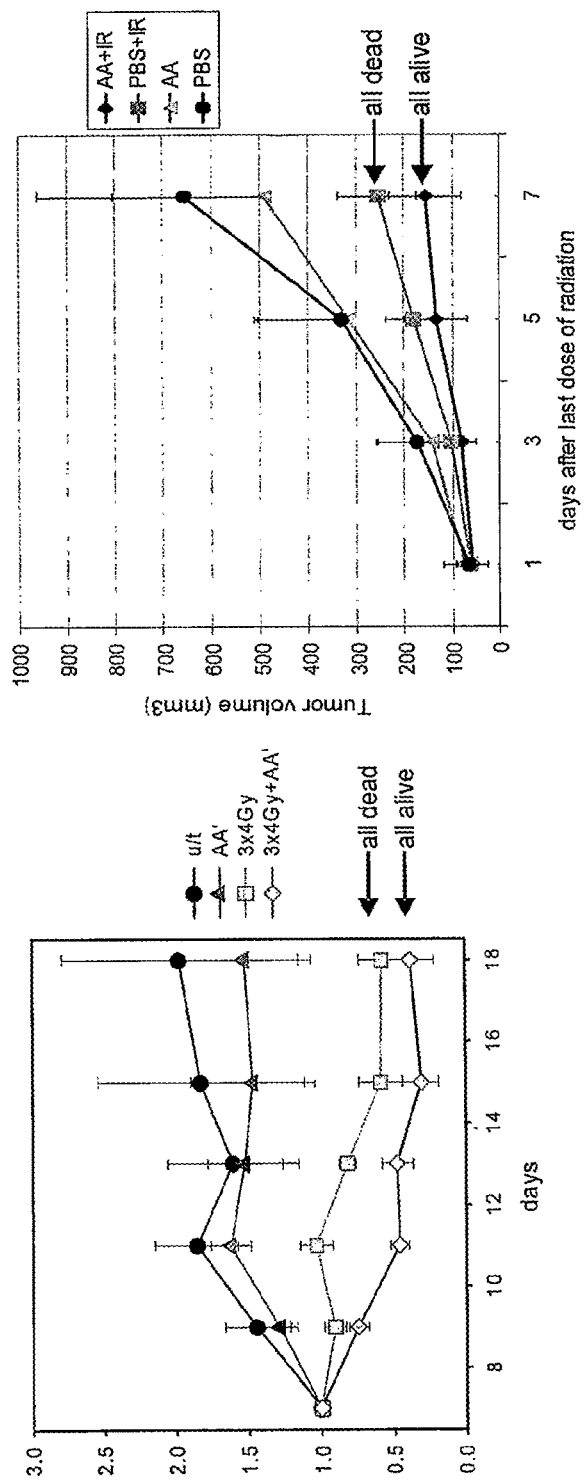


Fig. 23

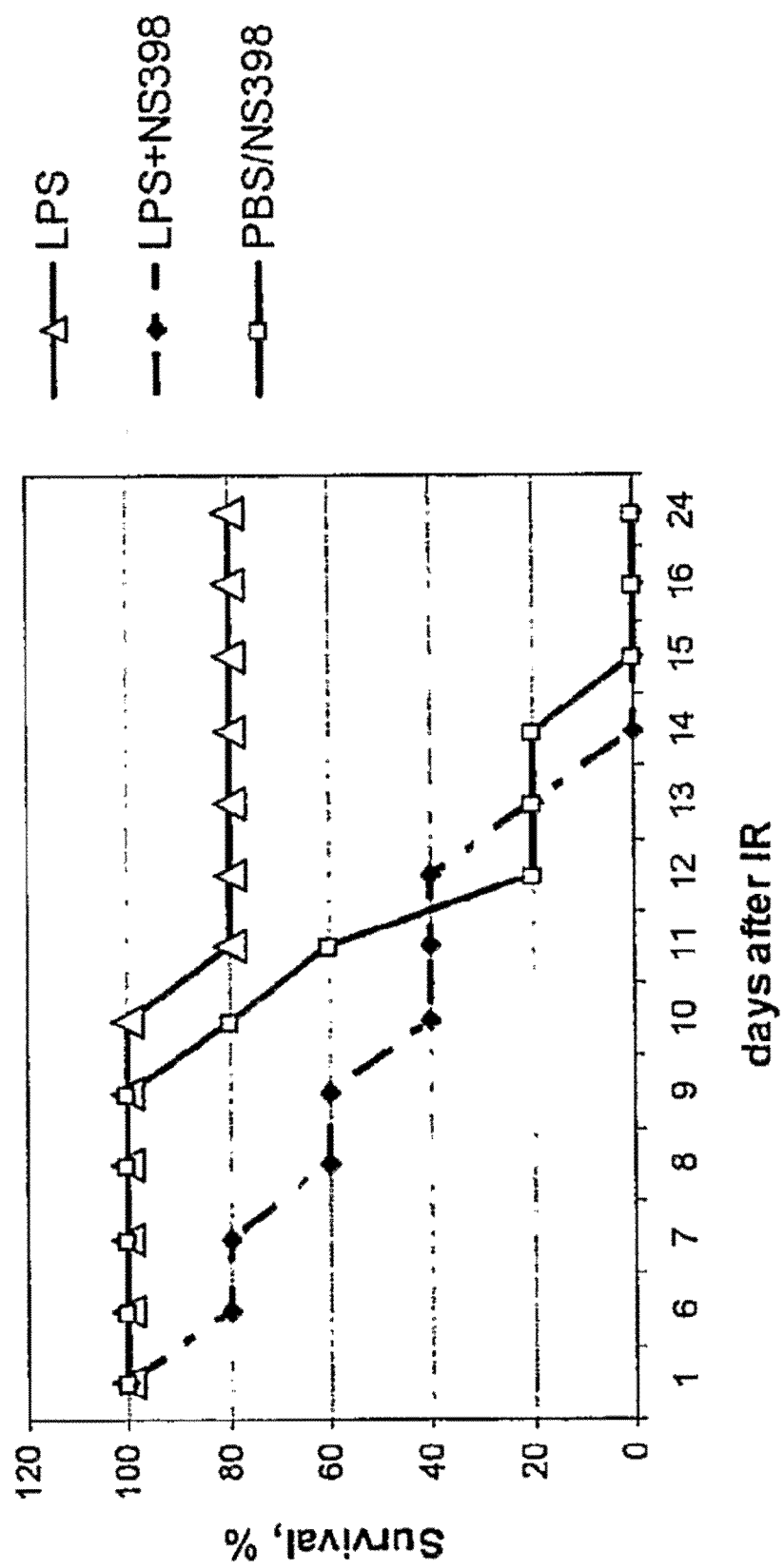


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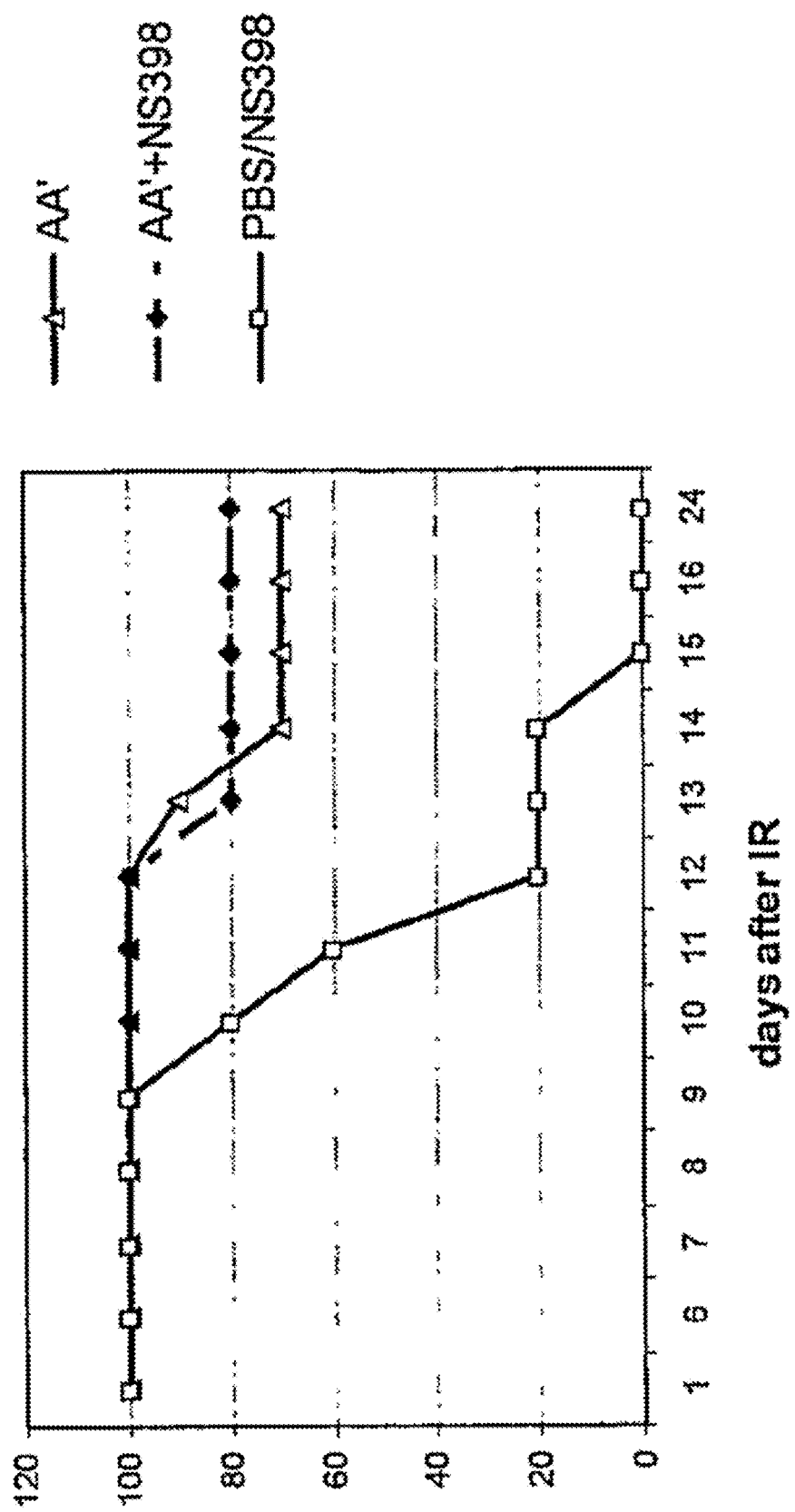


Fig. 24(A)

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Q5X5M6	1 MAQVINTINVASLCTAQEPNLGVSCRMQMTSIQRLSSGLRINSAKDDAAGLATSQRMTAQIRGMNQAVRNAND
Q6VMV6	1 MAQVINTNSLSLLTONNLNSQSSLGSAIERLSSGLRINSAKDDAAGQAIAANRFTANIKGLTQASRNAND
P13713	1 MAQVINTNSLSLMAQNNLNKSOSSLGTAIERLSSGLRINSAKDDAAGQAISNRFTANIKGLTQASRNAND
Q93RK8	1 --MRINHNIAALNTSRQLNAGSNSAAKONMEKLSSGLRINRAGDDAAGLAISENMRSQLRGLEMASKNAQD
Q02551	1 --MKVNTNISLRLTQEYLKRKNMNEGMLTQAERLSSGLRINSAKDDAAGLAVVTRMNVKSTGLDAASKNNSM
Q90912	1 MAQVINTNSLSLLTONNLNSQSSLSAIEERLSSGLRINSAKDDAAGQAIAANRFTANIKGLTQASRNAND
Q8GNT8	1 MAQVINTNSLSLMAQNNLNKSOSSLGTAIERLSSGLRINSAKDDAAGQAISNRFTANINGLTQASRNAND
Q9FAE7	1 MASTINTNVSSLTVAQRNLGSQSSLNTB1QRLLSSGLRINSAKDDAAGLAISENRTFSQIRGLNQAVRNAND
Q8ZP76	1 MA-VINTNSLSLLTONNLNSQSSLGTAIERLSSGLRINSAKDDAAGQAIAANRFTANVKGLTQARPNAND
Q7N5J4	1 MAQVINTNSLSLMAQNNLNRSQGTLLGSAIERLSSGLRINSAKDDAAGQAIAANRFTANVRGLTQAPRNAND
Q33578	1 -MTTINTNIGATAAQANMTKVNDQPNTAMTRLSTGLRINAACKDDAAGMAIGEKMTAQVGLNQAIRNAQD
Q56826	1 MASVINTNDGALLAQNNLTKXGILGSALERLSSGLRINSAKDDAAGQAIAANRFTANVKGLTQARPNAND
P42273	1 MAQVINTNLYSLVUTQNNLNRSQSSLGNAIERLSSGLRINSAKDDAAGQAIAANRFTSNINGLTQASRNAND
Q31059	1 --MVVQHNMQAAANSRMLGITTGDQSKESTEKLSSGPKINRAADAGLISBEKMRKQIRCLDQASTN ASD
Q7VZC2	1 MAAVINTNLYSLVVAQNNLNKBSQSSLGSAIERLSSGLRINSAKDDAAGQAIAANRFTANVKGLTQARPNAND
Q9F4A4	1 --MIINHNMALNNAHRNMGMNITAGKSMYKLISSGHLRINRAGDDAAGLAISEKMRGGQIRGLDQASHRNAQD
Q8P9C4	1 MAQVINTNMSLNAQRLNNTSSMALSICQLSSGKRTISASVDAAGLAISENRTFTQIRGLDVAARNAND
Q82UA3	1 MFQVINTNIALSLNAQRLNNTSRLSTALQRLLSSGLRINSAKDDAAGLAISENRTSQIRGMNQARPNAND
Q84IC5	1 -GFRINTNGASLNAQVNAGLMSRNLDSLARISSGLRINSAAADDASGLAIADSINTQANSILQQAINNAND
	;; * : ; : * ; : * ; : * ; : ; ; * ; :
Q53970	71 GISLAQTTEGALNEIMNLQVRELTVQGTSNSDSDLKSIQDEICCRLEEDIDRVSNQTFNGVKVLSD
P72151	71 GISLAQTAEGALNQSQSTNILQRLIKDIALLGANGNSNSDADRALQREBVAQQAELETRISDTTFFGGRKLLDG
Q5X5M6	71 GISLAQVAEGAMQETTINLQRLRELVSCAAMSTNNSSDRASIQSTISQLKSELERIAQNTTEFNGQRILDG
Q6VMV6	71 GISVAQTTEGALNEIMNLQVRELTVQGTSNSDSDLSSIQDEITQRLSEIDRVSSEQTQFNGVKVLAB
P13713	71 GISLAQTTEGALNEVNDNLQNTKRLTVQAGNSNSTSDLKSIQDEITQRLSEINRRISEQTQFNGVKVLSD
Q93RK8	69 GISLIQTTGECALNETSILORMSRELCAANDTNTPDRSELOKEMDQSLASERVTTRISTTTEFNTKLLDG
Q02551	71 GISLIQTTADEALSSMSSILQRMQLAVCSMGPSPSDEDRKQYTAEPGLSLIKELDHVADTTNNNKKLLDQ
Q99012	69 GISVAQTTEGALSEIMNLQPIRELVSCTATGTCNSDSDLNSIQDEITQRLSEIDRVSSEQTQFNGVKVLAS
Q8GNT8	71 GISLAQTTEGALNBVDNQNLQVRELTVQAGNSNSSSDLQSIQDEITQRLSEIDRISQQTDFNGVKVLSK
Q9FAE7	71 GISLAQTAEGALKSTGDIQHVRRELQVAGSANATNSSEDRKAIIQAEVGQLLSEMDRILAQNTEFNGQKLLDG
Q8ZP76	70 GISIAQTTGSLNEIMNLQVRELTVQAGNSNSSSDLQSIQDEITQRLSEIDRVSQDQTFNGVKVLAB
Q7N5J4	71 GISIAQTTTEGALNEIMNLQVRELTVQAGNSNSSESIDLKSIQDEITQRLSEIDRVSQDQTFNGVKVLRE
Q33578	70 GKNLVDTITGAHEVEVSSMQLPRLRELAVQSSMTNTADRGSLAAEKGQLIAEINRVAESTTUFNGMKVLLG
Q56826	71 GISIAQTTTEGALNEIMNLQVRELTVQAGNSNSDLESIQDEITQRLSEIDRISTQTQFNGIKVLNG
P42273	71 GSVSQTTTEGALNEIMNLQVRELTVQAGNSNSDLESIQDEITQRLSEIDRVSSEQTQFNGVKVLSG
Q31059	69 GISAVQTAEGALTEVHSMQLRMNELAQNAGTNSNSDRSSIQDEINQLTTEIDRAVETTKPNETYLLKG
Q7VZC2	71 GISIAQTTTEGALNEIMNLQVRELTVQAGNSNSDLESIQDEITQRLSEIDRVSQDQTFNGVKVLKS
Q9F4A4	69 GISLIQTAEGALAETHSISLQRMRELQVQ SANDINVAADRITAQDEITQRLSEIDRISGDTTEFNTQKLLDG
Q8P9C4	71 GISLAQTAEGALVEIGNNLQRELRELVSQSANATNSATDREALNSEVQQLTSEIDRVSANQTFNGVKLLNG
Q82UA3	71 GISLAQTAEGALVEMGNLQRELRELVSQSANATNSSEDDREALQKETQQLIDEIQRVGBQTSFNGVKLLDG
Q84IC5	70 ANSMLQTADEQLKILDTIKVKATDQACDQTAKTRAMIQGEINKLMEEIDNIAINTTYNGKQLLSD
	* : ; : * : ; : * : ; : * : ; : * : ; : * : ; : *

Fig. 24 (A) Continued

Q53970	141 DNO-MK--IQVGANDG-----	ETITIDLO-----	KID-VKSLG----LDGFN
P72151	141 SFGTTS--FQVGSNAY-----	ETIDISLQNASSASAIGSYQVG-SNGAGTVASVAGTA	
Q5X5M6	141 SFSGAS--FQVGANSN-----	QTINFSIG-----	SIK-ASSIGGIATATGTE
Q6VMV6	141 XNE-MK--IQVGANDG-----	ETITINLA-----	KID-AKTLG---LDGFN
P13713	141 DQK-LT--IQVGANDG-----	ETPDIDLK-----	KID-AKQLG---MDTF-
Q93RK8	139 TAONLT--FQIGANEG-----	QTMSLSIN-----	KMD-SE-----SLK
Q02551	139 TATGAATQVSIQASDKAN-----	DLENIDLFAKGGLSAGTTITLGSGSTVAGYSAALSVA	
Q09012	141 DQT-MK--IQVGANDG-----	ETIBELALD-----	KID-AKTLG---LDNPS
Q83NT8	141 DQK-LT--IQVGANDG-----	ETIDIDLK-----	NIN-AQSLG---LDKFN
Q9FAE7	141 SFGSAT--FQVGANAN-----	QTITATTGNFRINNY-GAQLT-ASASG--AATSGAS	
Q8ZF76	140 NTT-MS--IQVGANDG-----	ETIDINLQ-----	KID-SKSLG---LGSSYS
Q7N5J4	141 DSK-MT--IQVGANDN-----	EVIDIDLK-----	KID-KEALN---LGKFT
O33578	140 SFTGKQ--LQIGADSG-----	QTMAINVDSAATDIGAHKISSASTVVADAALTDIT	
Q56826	141 DVTEMK--IQVGANDN-----	ETIGIKLG-----	KIN-SEKLN---LKEFS
P42273	141 BKSKMT--IQVGTNDN-----	EVIRENLD-----	KID-NDTLG---VASDK
O31059	139 GNGDRT--VRVYAHAGLVGGLSQNTTRATFQMRLHEIGDSYTIGGITYKIG-AFTVK--EAMTALK		
Q7VZC2	141 MATDMTISIQVGAKDN-----	ETIDIKID-----	RNS-NWNLY---DAVGT
Q9F4A4	139 GFKG-K--FQIGANSN-----	QTVKLIDIG-----	NMS-AA-----SLG
Q8P9C4	141 DFSGAL--FQVGADAG-----	QTIGTINS-----	IVDAN-VDSLGS-KANFAAS
Q82UA3	141 SFASQI--FQVGANEG-----	ETIDETD-----	
Q84IC5	140 SFNSAQ--FQIGDKAN-----	QTVNATIG-----	STN-SAKVGQTRFETGAV
:			
:			

Fig. 24 (B)

Q53970	4.18	PLASIDSALKYDVRSSLGAIQRFDGATNLGNTVNLNSARSRIEDADAYTEVNMSKAQIILQQAGTSVLAQANQVQPNVLSLR-
P72151	4.01	AIAVVDNALAAIDAFRADIGAVONRFPTNLLNINISENTAARSRIKDTDEAETAALSKNOYLQOAGTALLAQANQLPQAVALSLLR--
Q5X5M6	2.87	AIRIDAALNSTVSNRANGALQNRFESTIMLQNSDNLSAARSRIODADYAAEMA SULTKNOYLQOAGTAMLAQANSLPOSVSLILGR--
Q6VMV6	4.00	FLETIDKALAKYDNRSLDLGAVQNRFDSAITNLGNTVNLSSARSRIEDADAYTEVNMSRAQIILQQAGTSVLAQANQVQPNVLSLR-
P13713	2.64	PLATDKALAQVDCRSLLGAVQNRFDSVIMLNSTVNLSSAQSRSRIOADADYATEVNMSRANILQQAGTSVLAQANQVQPNVLSLR-
Q93RK8	2.45	ALTIKIALDTSSERAKLGAVONRLEHTIMLGTSSSENTSAESRIRDDMASEMMEYTKNNILTOASOANTLAQANQOOPQVLQQLKG-
Q02551	4.81	VIGLADAALTKIMQRADMGAYNRLLETTAGLMGAYENMOASESBRIRDAEAEVSELTKQILVOSGTTAMLAQANMKEPNVSLKLQQI
Q90912	4.37	PLSKLDEALAKYDPLRSSLGAVQNRFDSAINTNLGNTVNLSSARSRIEDADAYTEVNMSRAQIILQQAGTSVLAQANQVQPNVLSLR-
Q8GNT8	3.29	PLATDKALISQVDELRSSLGAVQNRFDSVIMLNSTVNLSSAQSRSRIOADADYATEVNMSRANILQQAGTSVLAQANQVQPNVLSLR-
Q9FAE7	4.05	ALKIDIAALSAYNGRASEGALOSRFFETTYNLLOSTSEMSAQRASRIOADADYATEVNMSRANILQQAGTSVLAQANQVQPNVLSLR-
Q8ZF76	2.82	PLETHDDAIIKQDVDRSSLGAVQNRFESAINTNLNNTVTNLTSARSRIEDADAYTEVNMSRANILQQAGTSVLAQANQVQPNVLSLR-
Q7N5J4	2.68	PLETDLSALA0VDSRSSELGAIQRNLQESTVNLLNTVNLTSARSRIEDADAYTEVNMSRANILQQAGTSVLAQANQVQPNVLSLR-
Q33578	4.05	AIGVILVAISKISRSRSBGAWSRNLDTSTNUTISVQASQYNDADPAESTNLARSQIISQASTAMLAQANMSEKQNVLSLR-
Q56826	2.26	PLDTLDKALAQVDELRSSLGAVQNRFESTIMLNNTVNLSSAQSRSRIOADADYATEVNMSRANILQQAGTSVLAQANQVQPNVLSLR--
P42273	2.80	ALATIDNAISKVIDEBSRSKLGAIQRNLQESTVNLLNTVNLSSAQSRSRIOADADYATEVNMSRANILQQAGTSVLAQANQVQPNVLSLR--
Q31059	3.85	AIDASDAAKYSQISRSBGAWSRNLDTSTNUTISVQASQYNDADPAESTNLARSQIISQASTAMLAQANMSEKQNVLSLR-
Q7VZC2	3.04	ALSKDIDAKAVEQRSISLGAIQRNLQESTVNLLNTVNLSSAQSRSRIOADADYATEVNMSRANILQQAGTSVLAQANQVQPNVLSLR--
Q9F4A4	3.26	SIKTMNSAQSRSKLGAVQNRFESTIMLNNTSSENLTAAEGRYDUDMAKEMMASKNLISQOAAQMLGQANQOOPQVLQQLR--
Q8P9C4	3.12	ALEIVDKALTSVNSRSLAQSRSRIZADTDAEELTRTQILQQAGTAMLAQAKSTYSPONVLSLQQ
Q82U3	1.92	DDAKIVNSRSLAQSRSRIZADTAAELTRAQIILQQAGVAMSQANALDPNVLSLRR--
Q84TC5	4.03	VMDIEATALEIFIRANIGATQNCITSTTINLISVTVQVNKAESQIIRUVDFASESANYSKANLIAQSGSYAMAQANAQSQVNVLRLQ--

Fig. 25A

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AJ605785	1050	386	1.00E-107	348	215	51	AY353403	1518	196	2.00E-49	227	126	55
AJ536600	1050	385	1.00E-106	348	214	51	AY353402	1518	336	1.00E-91	178	177	99
AY249989	1050	384	1.00E-106	348	213	51	AY353401	1518	194	5.00E-49	106	104	98
AJ605764	1050	384	1.00E-106	348	213	51	AY353400	1518	336	1.00E-91	178	177	99
AB028472	1433	384	1.00E-106	348	213	51	AY353401	1518	194	5.00E-49	106	104	98
AJ605766	1050	383	1.00E-105	348	214	51	AY353400	1518	194	5.00E-49	106	104	98
AJ515904	1243	382	1.00E-105	348	213	51	AY353400	1518	194	5.00E-49	106	104	98
AY743588	829	378	1.00E-104	284	201	70	AY353399	1518	336	1.00E-91	178	177	99
X91047	2820	369	1.00E-101	313	206	55	AY353399	1518	194	5.00E-49	106	104	98
D32256	1240	367	1.00E-101	356	214	60	AY353398	1518	336	1.00E-91	178	177	99
BX571865	348813	364	1.00E-100	355	207	58	AY353398	1518	194	5.00E-49	106	104	98
AF221596	7867	356	8.00E-98	367	213	58	AY353397	1515	336	1.00E-91	178	177	99
AF221598	7867	355	1.00E-97	365	211	57	AY353397	1515	193	1.00E-48	106	103	97
AY649742	1518	336	1.00E-91	178	177	99	AY353396	1515	336	1.00E-91	178	177	99
AY649742	1518	194	5.00E-49	106	104	98	AY353396	1515	193	1.00E-48	105	103	97
AY649741	1518	336	1.00E-91	178	177	99	AY353395	1518	336	1.00E-91	178	177	99
AY649741	1518	194	5.00E-49	106	104	98	AY253395	1518	194	5.00E-49	106	104	98
AY649715	1518	336	1.00E-91	178	177	99	AY353394	1518	335	1.00E-91	178	177	99
AY649715	1518	197	5.00E-50	227	126	55	AY353394	1518	194	5.00E-49	106	104	98
AY649714	1518	336	1.00E-91	178	177	99	AY353393	1518	335	1.00E-91	178	177	99
AY649714	1518	195	2.00E-49	227	125	55	AY353393	1518	194	5.00E-49	106	104	98
AY649713	1518	335	1.00E-91	178	177	99	AY353392	1518	336	1.00E-91	178	177	99
AY649713	1518	195	2.00E-49	227	126	55	AY353392	1518	194	5.00E-49	106	104	98
AY649712	1518	336	1.00E-91	178	177	99	AY353391	1518	336	1.00E-91	178	177	99
AY649712	1518	194	5.00E-49	106	104	98	AY353391	1518	194	5.00E-49	106	104	98
AY649711	1518	336	1.00E-91	178	177	99	AY353389	1518	336	1.00E-91	178	177	99
AY649711	1518	194	5.00E-49	106	104	98	AY353389	1518	194	5.00E-49	106	104	98
AY649710	1518	336	1.00E-91	178	177	99	AY353388	1518	336	1.00E-91	178	177	99
AY649710	1518	194	5.00E-49	106	104	98	AY353388	1518	194	5.00E-49	106	104	98
AY649709	1518	336	1.00E-91	178	177	99	AY353387	1618	336	1.00E-91	178	177	99
AY649709	1518	194	5.00E-49	106	104	98	AY353387	1618	194	5.00E-49	106	104	98
AY649708	1618	336	1.00E-91	178	177	99	AY353386	1518	336	1.00E-91	178	177	99
AY649708	1618	194	5.00E-49	106	104	98	AY353386	1518	194	5.00E-49	106	104	95
AY649707	1518	336	1.00E-91	178	177	99	AY353385	1518	336	1.00E-91	178	177	99
AY649707	1518	197	8.00E-50	227	126	55	AY353385	1618	197	8.00E-50	227	126	55
AY649706	1518	336	1.00E-91	178	177	99	AY353384	1518	336	1.00E-91	178	177	99
AY649706	1518	194	4.00E-49	227	125	55	AY353384	1518	197	8.00E-50	227	126	55
AY649705	1518	336	1.00E-91	178	177	99	AY353383	1518	336	1.00E-91	178	177	99
AY649705	1518	197	8.00E-50	227	126	55	AY353383	1518	197	8.00E-50	227	126	55
AY353354	1518	336	1.00E-91	178	177	99	AY353381	1518	336	1.00E-91	178	177	99
AY353354	1518	194	4.00E-49	227	125	55	AY353381	1518	197	8.00E-50	227	126	55
AY353353	1527	336	1.00E-91	178	177	99	AY353380	1518	336	1.00E-91	178	177	99
AY353353	1527	197	5.00E-50	324	142	43	AY353380	1618	197	8.00E-50	227	126	55
AY353352	1527	336	1.00E-91	178	177	99	D78639	1630	336	1.00E-91	178	177	99
AY353352	1527	197	5.00E-50	324	142	43	D78639	1530	194	5.00E-49	106	104	98
AY353346	1515	336	1.00E-91	178	177	99	M84980	1518	336	1.00E-91	178	177	99
AY353346	1515	194	4.00E-49	309	141	45	M84980	1618	194	5.00E-49	106	104	98
AY353345	1515	336	1.00E-91	178	177	99	M84979	1518	336	1.00E-91	178	177	99
AY353345	1515	194	4.00E-49	309	141	45	M84979	1618	194	5.00E-49	106	104	98
AY353342	1527	336	1.00E-91	178	177	99	M84972	1518	336	1.00E-91	178	177	99
AY353342	1527	194	5.00E-49	106	104	98	M84972	1618	194	5.00E-49	106	104	98
AY353341	1527	336	1.00E-91	178	177	99	M84976	1518	336	1.00E-91	178	177	99
AY353341	1527	194	5.00E-49	106	104	98	M84976	1618	194	5.00E-49	106	104	98
AY353341	1527	336	1.00E-91	178	177	99	M84976	1518	336	1.00E-91	178	177	99
AY353340	1527	194	5.00E-49	106	104	98	M84976	1618	194	5.00E-49	106	104	98
AY353340	1527	336	1.00E-91	178	177	99	M84975	1518	336	1.00E-91	178	177	99
AY353340	1527	194	5.00E-49	106	104	98	M84975	1618	194	5.00E-49	106	104	98
AY353340	1527	336	1.00E-91	178	177	99	M84973	1518	336	1.00E-91	178	177	99
AY353340	1527	194	5.00E-49	105	104	98	M84973	1618	194	5.00E-49	106	104	98
AY353340	1518	336	1.00E-91	178	177	99	U06455	1867	336	1.00E-91	178	177	99
AY353340	1518	194	4.00E-49	227	125	55	U06455	1867	194	5.00E-49	106	104	98
AY353340	1518	336	1.00E-91	178	177	99	U06227	1876	336	1.00E-91	178	177	99
AY353340	1518	194	4.00E-49	106	104	98	U06227	1876	197	5.00E-50	324	142	43
AY353340	1518	336	1.00E-91	178	177	99	U05225	1867	336	1.00E-91	178	177	99
AY353340	1518	197	5.00E-50	227	126	55	U05225	1867	197	8.00E-50	227	126	55
AY353340	1518	336	1.00E-91	178	177	99	U06205	1864	336	1.00E-91	178	177	99
AY353340	1518	197	5.00E-50	227	126	55	U06205	1864	193	1.00E-48	106	103	97
AY353340	1518	336	1.00E-91	178	177	99	U06204	1876	336	1.00E-91	178	177	99
AY353340	1518	194	5.00E-49	106	104	98	U06204	1876	194	4.00E-49	227	125	55
AY353340	1518	336	1.00E-91	178	177	99	U06203	1867	336	1.00E-91	178	177	99

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
U06203	1867	197	5.00E-50	227	126	55	M84978	1518	192	1.00E-48	106	103	97
U06202	1876	336	1.00E-91	178	177	99	M84977	1518	333	5.00E-91	178	176	98
U06202	1876	194	5.00E-49	106	104	98	M84977	1518	194	5.00E-49	106	104	98
U06201	1876	336	1.00E-91	178	177	99	M27219	1544	333	7.00E-91	354	196	55
U06201	1876	194	5.00E-49	106	104	98	U05298	1398	330	5.00E-90	177	175	98
U06200	1876	336	1.00E-91	178	177	99	U05298	1398	171	6.00E-42	91	91	100
U06200	1876	197	8.00E-50	305	140	45	AY353413	1515	330	6.00E-90	178	173	97
U06197	1867	336	1.00E-91	178	177	99	AY353413	1515	187	6.00E-47	227	120	52
U06197	1867	194	4.00E-49	227	125	55	U06199	1864	330	6.00E-90	178	173	97
M84974	1867	336	1.00E-91	178	177	99	U06199	1864	187	6.00E-47	227	120	52
M84974	1867	194	5.00E-49	106	104	98	U05301	1512	328	2.00E-89	177	172	97
AY353535	1518	335	2.00E-91	178	176	98	U05301	1512	187	6.00E-47	227	120	52
AY353535	1518	194	4.00E-49	227	125	55	BX950851	5064019	327	5.00E-89	302	187	61
U06206	1876	335	2.00E-91	178	176	98	BX950851	5064019	131	4.00E-30	126	71	56
U06206	1876	193	1.00E-48	106	103	97	BX950851	5064019	88.6	4.00E-17	152	63	41
AY353537	1515	334	3.00E-91	178	176	98	L33488	1282	319	1.00E-86	365	203	55
AY353537	1515	189	2.00E-47	106	101	95	AE017132	290155	318	3.00E-86	368	202	54
AY353536	1515	334	3.00E-91	178	176	98	AJ414150	335050	318	3.00E-86	368	202	54
AY353536	1515	190	7.00E-48	280	139	49	AE013850	10283	318	3.00E-86	368	202	54
AY353382	1518	334	3.00E-91	178	176	98	BX936398	4744671	98.2	5.00E-20	207	61	29
AY353382	1518	194	4.00E-49	227	125	55	BX936398	4744671	60.1	2.00E-08	115	37	32
AY649717	1515	334	4.00E-91	178	176	98	AY649717	1515	315	2.00E-85	368	201	54
AY649716	1515	191	6.00E-48	301	136	45	U12963	1484	315	3.00E-85	166	165	99
AY649716	1515	334	4.00E-91	178	176	98	U12963	1484	194	5.00E-49	106	104	98
Z15086	1515	334	4.00E-91	177	176	99	AY337465	1467	311	2.00E-84	263	178	67
Z15086	1515	196	2.00E-49	227	126	55	AY337465	1467	160	8.00E-39	103	83	80
Z15072	1515	334	4.00E-91	177	176	99	AY353477	1518	311	2.00E-84	244	175	71
Z15072	1515	197	5.00E-50	227	126	55	AY353477	1518	175	3.00E-43	140	96	68
Z15071	1515	334	4.00E-91	177	176	99	AY353476	1518	311	2.00E-84	244	175	71
Z15071	1515	191	5.00E-49	106	104	98	AY353476	1518	175	3.00E-43	140	96	68
Z15070	1524	334	4.00E-91	177	176	99	AY353475	1518	311	2.00E-84	244	175	71
Z15070	1524	194	5.00E-49	106	104	98	AY353475	1518	175	3.00E-43	140	96	68
Z15069	1515	334	4.00E-91	177	176	99	AY353474	1518	311	2.00E-84	244	175	71
Z15069	1515	194	5.00E-49	106	104	98	AY353474	1518	175	3.00E-43	140	96	68
Z15068	1515	334	4.00E-91	177	176	99	AB108532	3325	310	5.00E-84	273	180	65
Z15068	1515	194	5.00E-49	106	104	98	AB108532	3325	166	2.00E-40	280	121	43
Z15067	1515	334	4.00E-91	177	176	99	L33457	1300	310	8.00E-84	359	194	54
Z15067	1515	194	5.00E-49	106	104	98	AY337472	1467	309	1.00E-83	263	177	67
Z15066	1515	334	4.00E-91	177	176	99	AY337472	1467	160	8.00E-39	103	83	80
Z15066	1515	197	8.00E-50	227	126	55	AF345848	1464	308	2.00E-83	256	177	69
Z15065	1515	334	4.00E-91	177	176	99	AF345848	1464	156	4.00E-38	103	82	79
Z15065	1515	197	5.00E-50	227	126	55	AJ567919	1392	308	3.00E-83	262	173	66
Z15054	1524	334	4.00E-91	177	176	99	AJ567919	1392	117	6.00E-26	75	59	78
Z15054	1524	197	5.00E-50	324	142	43	AY353479	1518	308	3.00E-83	244	172	70
AY353414	1515	334	4.00E-91	178	176	98	AY353479	1518	171	5.00E-42	140	94	67
AY353414	1515	191	6.00E-48	301	136	45	AY353478	1518	308	3.00E-83	244	172	70
U05303	1515	334	4.00E-91	177	176	99	AY353478	1518	173	9.00E-43	140	95	67
U05303	1515	197	5.00E-50	227	126	55	AY337475	1578	306	9.00E-83	299	184	61
U05302	1524	334	4.00E-91	177	176	99	AY337475	1578	149	2.00E-35	92	76	82
U05302	1524	194	5.00E-49	106	104	98	AB128919	1996	306	9.00E-83	299	184	61
U05300	1515	334	4.00E-91	177	176	99	AB128919	1996	150	7.00E-36	92	77	83
U05300	1515	194	5.00E-49	106	104	98	AY337476	1578	305	2.00E-82	299	184	61
U05299	1515	334	4.00E-91	177	176	99	AY337476	1578	149	2.00E-35	92	76	82
U05299	1515	194	5.00E-49	106	104	98	U47614	1762	305	2.00E-82	274	170	62
U05297	1515	334	4.00E-91	177	176	99	U47614	1762	142	3.00E-33	115	74	64
U05297	1515	194	5.00E-49	106	104	98	AE005174	5528445	305	3.00E-82	274	169	61
U05296	1524	334	4.00E-91	177	176	99	AE005174	5528445	142	3.00E-33	115	74	64
U05296	1524	194	5.00E-49	105	104	98	AE005174	5528445	51.6	5.00E-06	299	66	22
U05295	1515	334	4.00E-91	177	176	99	AE005174	5528445	33.5	1.5	185	42	22
U05295	1515	196	1.00E-49	227	126	55	AY337468	1758	305	3.00E-82	274	169	61
U06226	1864	334	4.00E-91	178	176	98	AY337468	1758	142	3.00E-33	115	74	64
U06226	1864	191	6.00E-48	301	136	45	AY249992	1758	305	3.00E-82	267	168	62
U06198	1864	334	4.00E-91	178	176	98	AY249992	1758	142	3.00E-33	115	74	64
U06198	1864	191	6.00E-48	301	136	45	BA000007	5498450	305	3.00E-82	274	169	61
AY649696	1518	333	5.00E-91	178	176	98	BA000007	5498450	142	3.00E-33	115	74	64
AY649696	1518	194	5.00E-49	106	104	98	BA000007	5498450	51.6	5.00E-06	299	66	22
M84978	1518	333	5.00E-91	178	176	98	BA000007	5498450	33.5	1.5	185	42	22

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AF228496	1758	305	3.00E-82	274	169	61	AY250027	1818	296	7.00E-80	222	157	70
AF228496	1758	142	3.00E-33	115	74	64	AY250027	1818	139	2.00E-32	163	81	49
AF228494	1758	305	3.00E-82	267	168	62	AB028481	2168	296	7.00E-80	222	157	70
AF228494	1758	142	3.00E-33	115	74	64	AB028481	2168	139	2.00E-32	163	81	49
AF228493	1758	305	3.00E-82	267	168	62	AY250029	1311	296	1.00E-79	300	175	58
AF228493	1758	142	3.00E-33	115	74	64	AY250029	1311	135	4.00E-31	92	67	72
AF228492	1758	305	3.00E-82	267	168	62	AF345847	1497	296	1.00E-79	178	155	87
AF228492	1758	142	3.00E-33	115	74	64	AF345847	1497	155	2.00E-37	174	94	54
AF228491	1758	305	3.00E-82	267	168	62	AY380836	1714	296	1.00E-79	223	159	71
AF228491	1758	142	3.00E-33	115	74	64	AY380836	1714	140	7.00E-33	114	74	64
AF228490	1758	305	3.00E-82	267	168	62	AB028480	2259	296	1.00E-79	302	176	58
AF228490	1758	142	3.00E-33	115	74	64	AB028480	2259	133	1.00E-30	102	66	64
AF228489	1758	305	3.00E-82	274	169	61	L07389	1788	296	1.00E-79	223	156	69
AF228489	1758	142	3.00E-33	115	74	64	L07389	1788	135	3.00E-31	113	70	61
AF228488	1758	305	3.00E-82	274	169	61	AY250001	1655	295	1.00E-79	237	161	67
AF228488	1758	142	3.00E-33	115	74	64	AY250001	1655	141	5.00E-33	295	105	35
AF228487	1758	305	3.00E-82	274	169	61	AF194946	1654	295	1.00E-79	237	161	67
AF228487	1758	142	3.00E-33	115	74	64	AF194946	1654	139	2.00E-32	293	104	35
D16821	2006	305	3.00E-82	299	183	61	AY337480	1512	295	2.00E-79	178	155	87
D16821	2006	148	3.00E-35	92	76	82	AY337480	1512	158	4.00E-38	265	111	41
D16820	1851	305	3.00E-82	299	183	61	AY250016	1638	295	2.00E-79	218	157	72
D16820	1851	148	3.00E-35	92	76	82	AY250016	1638	140	7.00E-33	172	85	49
AB028474	2329	305	3.00E-82	267	168	62	Z36877	1925	295	2.00E-79	222	156	70
AB028474	2329	142	3.00E-33	115	74	64	Z36877	1925	133	1.00E-30	102	66	64
AY337477	1578	304	4.00E-82	299	183	61	AY337483	1674	295	2.00E-79	284	169	59
AY337477	1578	149	2.00E-35	92	76	82	AY337483	1674	141	5.00E-33	115	71	61
AF228495	1758	304	5.00E-82	274	169	61	AY250028	1344	295	2.00E-79	274	171	62
AF228495	1758	142	3.00E-33	115	74	64	AY250028	1344	134	6.00E-31	92	67	72
AY337478	1578	303	1.00E-81	299	183	61	AY250010	1740	295	2.00E-79	236	161	68
AY337478	1578	145	4.00E-34	92	74	80	AY250010	1740	140	9.00E-33	116	72	62
L07386	1755	303	1.00E-81	267	169	63	AY250008	1674	295	2.00E-79	284	169	59
L07386	1755	141	5.00E-33	115	73	63	AY250008	1674	142	3.00E-33	115	71	61
AB128917	8242	301	4.00E-81	296	186	62	AY249999	1689	295	2.00E-79	257	168	65
AB128917	8242	141	5.00E-33	87	75	86	AY249999	1689	142	2.00E-33	273	102	37
AY337481	1604	300	9.00E-81	177	157	88	AY249990	1311	295	3.00E-79	291	174	59
AY337481	1604	147	7.00E-35	92	75	81	AY249990	1311	135	4.00E-31	92	67	72
AY249991	1647	300	9.00E-81	266	171	64	AF345850	1638	295	3.00E-79	218	157	72
AY249991	1647	140	7.00E-33	102	69	67	AF345850	1638	140	7.00E-33	172	85	49
AY250013	1668	299	1.00E-80	295	171	57	AB028473	1867	295	3.00E-79	291	174	59
AY250013	1668	140	7.00E-33	116	71	61	AB028473	1867	135	4.00E-31	92	67	72
AY337474	1788	298	2.00E-80	223	157	70	AY337473	1344	294	4.00E-79	290	171	68
AY337474	1788	137	7.00E-32	113	71	62	AY337473	1344	137	7.00E-32	314	115	36
AY337471	1788	298	2.00E-80	223	157	70	AY250018	1344	294	4.00E-79	290	171	58
AY337471	1788	137	7.00E-32	113	71	62	AY250018	1344	137	1.00E-31	314	114	36
AY250023	1707	298	2.00E-80	223	160	71	AY249994	2013	294	4.00E-79	247	162	65
AY250023	1707	140	7.00E-33	114	74	64	AY249994	2013	147	9.00E-35	312	104	33
AY249997	1788	298	2.00E-80	223	157	70	AF079163	2078	294	4.00E-79	247	162	65
AY249997	1788	137	7.00E-32	113	71	62	AF079163	2078	147	9.00E-35	312	104	33
AE016762	311143	298	2.00E-80	223	157	70	AY250026	1695	294	5.00E-79	222	156	70
AE016762	311143	138	3.00E-32	255	106	41	AY250026	1695	133	1.00E-30	102	66	64
AB028477	2277	298	2.00E-80	223	160	71	AF345851	1695	294	5.00E-79	222	156	70
AB028477	2277	141	4.00E-33	270	104	38	AF345851	1695	132	2.00E-30	102	66	64
AB028475	2359	298	2.00E-80	223	157	70	AE0056744607203	293	6.00E-79	202	154	76	
AB028475	2359	137	7.00E-32	113	71	62	AE0056744607203	293	140	7.00E-33	116	71	61
AB028471	2351	298	2.00E-80	223	157	70	AY337485	1323	293	6.00E-79	288	177	61
AB028471	2351	138	3.00E-32	255	106	41	AY337485	1323	137	1.00E-31	312	109	34
L07387	1788	298	2.00E-80	223	157	70	AF543692	1497	293	6.00E-79	178	153	85
L07387	1788	139	2.00E-32	255	106	41	AF543692	1497	155	2.00E-37	174	94	54
AJ566339	1634	298	3.00E-80	246	163	66	AE016934	290582	293	6.00E-79	202	154	76
AJ566339	1634	89.7	2.00E-17	83	44	53	AE016934	290582	140	7.00E-33	116	71	61
AJ566338	1634	298	3.00E-80	246	163	66	AY250003	1731	293	6.00E-79	299	171	57
AJ566338	1634	89.7	2.00E-17	83	44	53	AY250003	1731	142	2.00E-33	287	111	38
AY250005	1767	297	6.00E-80	219	162	73	D16819	1966	293	6.00E-79	202	154	76
AY250005	1767	142	2.00E-33	149	83	55	D16819	1966	140	7.00E-33	116	71	61
AB028476	2338	297	6.00E-80	219	162	73	AY250024	1719	293	8.00E-79	229	160	69
AB028476	2338	142	2.00E-33	149	83	55	AY250024	1719	144	8.00E-34	116	73	62
AY337466	1344	296	7.00E-80	290	172	59	AY249998	1653	293	8.00E-79	202	154	76
AY337466	1344	135	2.00E-31	181	87	48	AY249998	1653	140	7.00E-33	116	71	61

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AF169321	1653	293	6.00E-79	202	154	76	AY649737	1281	288	3.00E-77	214	161	75
AF169321	1653	138	4.00E-32	276	102	36	AY649737	1281	150	7.00E-36	100	79	79
AY337489	1311	293	1.00E-78	291	169	58	AY649736	1269	288	3.00E-77	214	161	75
AY337489	1311	134	5.00E-31	92	67	72	AY649736	1269	148	3.00E-35	100	78	78
AY250020	1580	292	1.00E-78	179	149	83	AY649735	1275	288	3.00E-77	214	161	75
AY250020	1580	139	3.00E-32	113	69	61	AY649735	1275	150	7.00E-36	100	79	79
AY250012	1332	291	2.00E-78	288	176	61	AY649704	1509	288	3.00E-77	220	160	72
AY250012	1332	135	3.00E-31	315	107	33	AY649704	1509	157	7.00E-38	100	82	82
D26166	3294	291	2.00E-78	234	159	67	AY649703	1509	288	3.00E-77	220	160	72
D26166	3294	134	6.00E-31	100	66	66	AY649703	1509	157	7.00E-38	100	82	82
AF543693	1497	291	3.00E-78	178	152	86	AY250022	1506	288	3.00E-77	220	156	70
AF543693	1497	155	2.00E-37	174	94	54	AY250022	1506	134	8.00E-31	151	77	50
AY353496	1242	291	3.00E-78	275	175	63	AF169323	1506	288	3.00E-77	220	156	70
AY353496	1242	150	9.00E-36	100	79	79	AF169323	1506	134	8.00E-31	151	77	50
AY353486	1266	291	3.00E-78	276	170	61	AY353494	1269	288	3.00E-77	214	161	75
AY353486	1266	152	3.00E-36	102	79	77	AY353494	1269	148	3.00E-35	100	78	78
AY353485	1266	291	3.00E-78	276	170	61	AY353484	1269	288	3.00E-77	214	161	75
AY353485	1266	152	3.00E-36	102	79	77	AY353484	1269	150	7.00E-36	100	79	79
D26165	2520	291	3.00E-78	263	159	60	AY353444	1269	288	3.00E-77	214	161	75
D26165	2520	140	7.00E-33	114	74	64	AY353444	1269	148	3.00E-35	100	78	78
AY337479	1851	291	4.00E-78	191	151	79	AY353507	1269	288	3.00E-77	214	161	75
AY337479	1851	145	3.00E-34	287	109	37	AY353507	1269	150	7.00E-36	100	79	79
AY250014	1713	291	4.00E-78	179	150	83	AY353506	1269	288	3.00E-77	214	161	75
AY250014	1713	139	2.00E-32	234	88	37	AY353506	1269	150	7.00E-36	100	79	79
AY260002	1842	291	4.00E-78	191	151	79	AY353505	1281	288	3.00E-77	214	161	75
AY260002	1842	146	2.00E-34	265	109	38	AY353505	1281	151	5.00E-36	174	92	52
AJ567918	1359	291	4.00E-78	251	164	65	AY353503	1281	288	3.00E-77	214	161	75
AJ567918	1359	117	6.00E-26	75	69	78	AY353503	1281	149	1.00E-35	174	91	52
AJ271930	1713	291	4.00E-78	179	150	83	AY353499	1269	288	3.00E-77	214	161	75
AJ271930	1713	139	2.00E-32	234	88	37	AY353499	1269	150	7.00E-36	100	79	79
AB028478	2284	290	7.00E-78	229	159	68	AY353497	1275	288	3.00E-77	214	161	75
AB028478	2284	144	8.00E-34	116	73	62	AY353497	1275	150	7.00E-36	100	79	79
AB128916	11067	290	7.00E-78	296	177	59	AY353495	1269	288	3.00E-77	214	161	75
AB128916	11067	151	5.00E-36	293	112	38	AY353495	1269	150	7.00E-36	100	79	79
AY250019	1299	290	9.00E-78	286	169	59	AY353494	1269	288	3.00E-77	214	161	75
AY250019	1299	138	1.00E-31	253	102	40	AY353494	1269	150	7.00E-36	100	79	79
AY260015	1287	290	9.00E-78	267	156	62	AY353492	1269	288	3.00E-77	214	161	75
AY260015	1287	134	6.00E-31	127	74	58	AY353492	1269	150	7.00E-36	100	79	79
AF345849	1287	290	9.00E-78	267	156	62	AY353454	1506	288	3.00E-77	216	158	73
AF345849	1287	131	4.00E-30	127	73	57	AY353454	1506	156	1.00E-37	100	82	82
AY353488	1266	290	9.00E-78	271	170	62	AY353447	1503	288	3.00E-77	208	157	75
AY353488	1266	152	3.00E-36	102	79	77	AY353447	1503	156	1.00E-37	100	82	82
AY353487	1266	290	9.00E-78	271	170	62	AY353446	1503	288	3.00E-77	208	157	75
AY353487	1266	152	3.00E-36	102	79	77	AY353446	1503	156	1.00E-37	100	82	82
AY353483	1266	290	9.00E-78	175	153	87	AY353379	1509	288	3.00E-77	220	160	72
AY353484	1266	289	1.00E-77	175	153	87	AY353379	1509	157	7.00E-38	103	82	82
AY353484	1266	152	3.00E-36	102	79	77	AY353377	1509	288	3.00E-77	220	160	72
AY353482	1266	289	1.00E-77	175	153	87	AY353377	1509	157	7.00E-38	100	82	82
AY250017	1688	289	2.00E-77	233	162	69	AY353331	1503	288	3.00E-77	208	157	75
AY250017	1688	150	9.00E-36	279	102	36	AY353331	1503	154	5.00E-37	100	81	81
AY353457	1269	288	2.00E-77	284	175	61	AY353455	1506	287	4.00E-77	216	158	73
AY353457	1269	150	7.00E-36	100	79	79	AY353455	1506	156	1.00E-37	100	82	82
AY353546	1269	289	2.00E-77	284	175	61	AY337470	1281	287	6.00E-77	258	163	63
AY353546	1269	150	7.00E-36	100	79	79	AY337470	1281	138	4.00E-32	232	93	40
AY363645	1269	289	2.00E-77	284	175	61	AY649725	1503	287	6.00E-77	206	157	76
AY363645	1269	150	7.00E-36	100	79	79	AY649725	1503	152	2.00E-36	98	79	80
AY353332	1503	288	2.00E-77	208	157	75	AY649724	1503	287	6.00E-77	198	154	77
AY353332	1503	155	3.00E-37	100	81	81	AY649724	1503	156	1.00E-37	100	82	82
AY250021	1281	288	3.00E-77	258	164	63	AY649723	1503	287	6.00E-77	198	154	77
AY250021	1281	138	4.00E-32	232	93	40	AY649723	1503	156	1.00E-37	100	82	82
AY250011	1713	288	3.00E-77	179	149	83	AY250025	1497	287	6.00E-77	177	146	82
AY250011	1713	139	3.00E-32	237	91	38	AY250025	1497	149	2.00E-35	151	80	52
AF169322	1281	288	3.00E-77	258	164	63	X17440	2526	287	6.00E-77	177	146	82
AF169322	1281	136	2.00E-31	92	89	75	X17440	2526	151	4.00E-36	244	101	41
AY353498	1269	288	3.00E-77	237	164	69	U00096 4639675	287	6.00E-77	177	146	82	
AY353498	1269	148	3.00E-35	100	78	78	U00096 4639675	149	2.00E-35	151	80	52	
AY353493	1269	288	3.00E-77	237	164	69	U00096 4639675	33.1	2	147	37	25	
AY353493	1269	148	3.00E-35	100	78	78	AY353453	1269	287	6.00E-77	211	159	75

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AY353543	1269	150	7.00E-36	100	79	79	AJ566340	1223	85.1	4.00E-16	284	81	28
AY353457	1482	287	6.00E-77	198	154	77	AY649698	1488	285	3.00E-76	306	176	57
AY353457	1482	156	1.00E-37	100	82	82	AY649698	1488	181	4.00E-36	96	78	81
AY353458	1446	287	6.00E-77	245	162	66	AY249995	1263	137	1.00E-31	138	78	56
AY353456	1446	156	1.00E-37	100	82	82	AY249995	1263	285	3.00E-76	189	147	77
AY353453	1506	287	6.00E-77	206	157	76	AY657001	1488	285	3.00E-76	306	176	57
AY353453	1506	156	1.00E-37	100	82	82	AY657001	1488	151	4.00E-36	96	78	81
AY353452	1506	287	6.00E-77	206	157	76	BX321861	298050	285	3.00E-76	297	159	53
AY353452	1506	154	5.00E-37	100	81	81	AY353508	1269	285	3.00E-76	211	158	74
AY353451	1503	287	6.00E-77	198	154	77	AY353508	1259	150	7.00E-36	100	79	79
AY353451	1503	156	1.00E-37	100	82	82	AY353363	1488	285	3.00E-76	306	176	57
AY353450	1503	287	6.00E-77	206	157	76	AY353363	1488	151	4.00E-36	96	78	81
AY353450	1503	156	1.00E-37	100	82	82	AY353362	1488	285	3.00E-76	306	176	57
AY353449	1503	287	6.00E-77	206	157	76	AY353362	1488	151	4.00E-36	96	78	81
AY353449	1503	152	2.00E-36	98	79	80	AY353361	1488	285	3.00E-76	306	175	57
AY353448	1503	287	6.00E-77	198	154	77	AY353361	1488	151	4.00E-36	96	78	81
AY353448	1503	156	1.00E-37	100	82	82	AY353359	1488	285	3.00E-76	306	176	57
AY353444	1503	287	6.00E-77	198	154	77	AY353359	1488	151	4.00E-36	96	78	81
AY353444	1503	156	1.00E-37	100	82	82	AY353358	1488	285	3.00E-76	306	176	57
AY353443	1503	287	6.00E-77	198	154	77	AY353358	1488	151	4.00E-36	96	78	81
AY353443	1503	156	1.00E-37	100	82	82	AY353303	1506	285	3.00E-76	175	150	85
AY353378	1509	287	6.00E-77	220	160	72	AY363303	1506	155	3.00E-37	100	81	81
AY353378	1509	157	7.00E-38	100	82	82	AY353517	1506	284	4.00E-76	215	157	73
D90833	15605	287	6.00E-77	177	146	82	AY353517	1506	152	2.00E-36	98	79	80
D90833	15605	149	2.00E-35	151	80	52	AY353516	1506	284	4.00E-76	215	157	73
AB028479	2050	287	6.00E-77	177	146	82	AY353516	1506	152	2.00E-36	98	79	80
AB028479	2050	149	2.00E-35	151	80	52	AY353504	1281	284	4.00E-76	214	160	74
M14358	1667	287	6.00E-77	177	146	82	AY353504	1281	149	1.00E-35	174	91	52
M14358	1667	149	2.00E-35	151	80	52	AY353490	1500	284	4.00E-76	192	152	79
AY353509	1266	286	8.00E-77	175	152	86	AY353490	1500	153	1.00E-36	100	80	80
AY353509	1266	150	9.00E-36	100	79	79	AY353489	1500	284	4.00E-76	192	152	79
AY353501	1266	286	8.00E-77	175	152	86	AY353489	1500	153	1.00E-36	100	80	80
AY353501	1266	150	9.00E-36	100	79	79	AY374782	1263	284	5.00E-76	189	147	77
AY353502	1266	286	8.00E-77	175	152	86	AY374782	1263	135	4.00E-31	138	77	55
AY353650	1266	150	9.00E-36	100	79	79	AY649732	1521	284	5.00E-76	175	150	85
AY353445	1503	286	8.00E-77	198	154	77	AY649732	1521	149	2.00E-35	91	76	83
AY353445	1503	152	2.00E-36	98	79	80	AY649701	1521	284	5.00E-76	175	150	85
AY374787	1281	286	1.00E-76	258	163	63	AY649701	1521	149	2.00E-35	91	76	83
AY374787	1281	136	1.00E-31	232	92	39	AY649700	1521	284	5.00E-76	175	150	85
AY353334	1508	286	1.00E-76	206	157	76	AY649700	1521	150	9.00E-36	91	77	84
AY353334	1506	156	1.00E-37	100	82	82	AE016837	300247	284	5.00E-76	175	150	85
AY353333	1506	286	1.00E-76	206	157	76	AE016837	300247	149	2.00E-35	91	76	83
AY353333	1506	156	1.00E-37	100	82	82	X03395	1530	284	6.00E-76	175	150	85
AB128918	12979	286	1.00E-76	291	170	68	X03395	1530	149	2.00E-35	91	76	83
AB128918	12979	146	2.00E-34	95	75	78	AL627272	245050	284	5.00E-76	175	150	85
AY649702	1509	286	1.00E-76	220	169	72	AL627272	245050	149	2.00E-35	91	76	83
AY649702	1509	157	7.00E-38	100	82	82	AY353434	1260	284	5.00E-76	175	150	85
AY353350	1500	286	1.00E-76	209	158	75	AY353434	1260	149	2.00E-35	91	76	83
AY353350	1500	158	3.00E-38	338	127	37	AY353433	1260	284	5.00E-76	175	150	85
AY353349	1500	286	1.00E-76	209	158	75	AY353433	1260	149	2.00E-35	91	76	83
AY353349	1500	156	4.00E-38	338	127	37	AY353378	1521	284	5.00E-76	175	150	85
AY353348	1500	286	1.00E-76	209	158	75	AY363376	1521	150	9.00E-36	91	77	84
AY353348	1500	158	3.00E-38	338	127	37	AY353375	1521	284	5.00E-76	175	150	85
AY353347	1500	286	1.00E-76	209	158	75	AY353375	1521	149	2.00E-35	91	76	83
AY353347	1500	158	4.00E-38	338	127	37	AY353375	1521	284	5.00E-76	175	150	85
AY353346	1500	286	1.00E-76	209	158	75	AY353374	1521	149	2.00E-35	91	76	83
AY353346	1500	157	5.00E-38	338	127	37	AY353373	1521	284	5.00E-76	175	150	85
AY353345	1500	286	1.00E-76	209	158	75	AY353373	1521	149	2.00E-35	91	76	83
AY353345	1500	158	4.00E-38	338	127	37	AY653372	1521	284	5.00E-75	175	150	85
AY353345	1500	286	1.00E-76	209	158	75	AY353372	1521	149	2.00E-35	91	76	83
AF169320	1263	285	2.00E-34	269	160	59	AY353371	1521	284	5.00E-76	175	150	85
AF169320	1263	135	4.00E-31	138	77	55	AY353371	1521	149	2.00E-35	91	76	83
AY353502	1266	285	2.00E-76	175	151	88	AY353371	1521	149	2.00E-35	91	76	83
AY353502	1266	150	9.00E-36	100	79	79	AY353302	1521	284	5.00E-76	175	150	85
AY353360	1488	285	2.00E-76	306	176	57	AY353302	1521	150	9.00E-36	91	77	84
AY353360	1488	149	2.00E-35	89	76	85	L21912	2001	284	5.00E-76	175	150	85
AJ566341	1223	285	3.00E-76	179	145	81	L21912	2001	149	2.00E-35	91	76	83
AJ566341	1223	85.1	4.00E-15	284	81	28	CP0000264585229	283	6.00E-76	279	165	59	
AJ566340	1223	285	3.00E-78	179	145	81	CP0000264585229	278	2.00E-74	175	147	84	

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
CP000264585229	156	1.00E-37	100	82	82		AY353275	1521	155	3.00E-37	100	81	81
CP000264585229	155	3.00E-37	100	81	81		U17172	1521	282	2.00E-75	232	159	68
AE008826	20513	283	6.00E-76	232	160	68	U17172	1521	155	3.00E-37	100	81	81
AE008826	20513	155	3.00E-37	100	81	81	AY353292	1521	281	2.00E-75	268	163	60
AY353521	1521	283	6.00E-76	232	160	68	AY353292	1521	155	3.00E-37	100	81	81
AY353521	1521	155	3.00E-37	100	81	81	AY353283	1521	281	2.00E-75	268	162	60
AY353271	1521	283	6.00E-76	232	160	68	AY353283	1521	155	3.00E-37	100	81	81
AY353271	1521	155	3.00E-37	100	81	81	AY353263	1521	281	2.00E-75	268	163	60
AY353269	1521	283	6.00E-76	232	160	68	AY353263	1521	155	3.00E-37	100	81	81
AY353269	1521	155	3.00E-37	100	81	81	U17175	1521	281	2.00E-75	228	156	68
AY353268	1521	283	6.00E-76	232	160	68	U17175	1521	155	3.00E-37	100	81	81
AY353268	1521	155	3.00E-37	100	81	81	AY353542	1485	281	3.00E-75	175	148	84
AY353267	1521	283	6.00E-76	232	160	68	AY353542	1485	152	2.00E-36	98	79	80
AY353267	1521	155	3.00E-37	100	81	81	AY353541	1485	281	3.00E-75	175	148	84
AY353266	1521	283	6.00E-76	232	160	68	AY353541	1485	152	2.00E-36	98	79	80
AY353266	1521	155	3.00E-37	100	81	81	AY353540	1485	281	3.00E-75	175	148	84
AY353264	1521	283	6.00E-76	232	160	68	AY353540	1485	152	2.00E-36	98	79	80
AY353264	1521	155	3.00E-37	100	81	81	AY353539	1488	281	3.00E-75	175	148	84
AY353259	1506	283	6.00E-76	295	169	57	AY353539	1488	148	4.00E-35	98	77	78
AY353259	1506	151	4.00E-36	98	78	79	AY353538	1488	281	3.00E-75	175	148	84
AF045151	1521	283	6.00E-76	232	160	68	AY353538	1488	152	2.00E-36	98	79	80
AF045151	1521	155	3.00E-37	100	81	81	AY353531	1521	281	3.00E-75	175	148	84
U17177	1521	283	6.00E-76	232	160	68	AY353531	1521	155	3.00E-37	100	81	81
U17177	1521	155	3.00E-37	100	81	81	AY353530	1521	281	3.00E-75	175	148	84
U17176	1521	283	6.00E-76	232	160	68	AY353530	1521	155	3.00E-37	100	81	81
U17176	1521	155	3.00E-37	100	81	81	AY353529	1521	281	3.00E-75	175	148	84
U17173	1521	283	6.00E-76	232	160	68	AY353529	1521	154	8.00E-37	100	80	80
U17173	1521	155	3.00E-37	100	81	81	AY353528	1521	281	3.00E-75	175	148	84
AY353272	1521	283	8.00E-76	232	160	68	AY353528	1521	155	3.00E-37	100	81	81
AY353272	1521	155	3.00E-37	100	81	81	AY353518	1488	281	3.00E-75	175	148	84
AJ243796	1368	283	1.00E-75	247	162	65	AY353518	1488	155	3.00E-37	245	105	42
AJ243796	1368	137	6.00E-32	87	70	80	AY353512	1485	281	3.00E-75	175	148	84
AJ243795	1368	283	1.00E-75	247	162	65	AY353512	1485	158	3.00E-38	272	108	39
AJ243795	1368	137	6.00E-32	87	70	80	AY353470	1500	281	3.00E-75	175	148	84
AY353525	1521	283	1.00E-75	229	160	69	AY353470	1500	154	8.00E-37	100	80	80
AY353525	1521	154	6.00E-37	150	87	58	AY353469	1500	281	3.00E-75	175	148	84
AY353524	1521	283	1.00E-75	229	160	69	AY353469	1500	154	8.00E-37	100	80	80
AY353524	1521	154	6.00E-37	150	87	58	AY353439	1485	281	3.00E-75	175	148	84
AY353523	1521	283	1.00E-75	229	160	69	AY353439	1485	152	2.00E-36	98	79	80
AY353523	1521	154	8.00E-37	100	80	80	AY353438	1485	281	3.00E-75	175	148	84
AY353522	1521	283	1.00E-75	229	160	69	AY353438	1485	152	2.00E-36	98	79	80
AY353522	1521	154	8.00E-37	100	80	80	AY353436	1488	281	3.00E-75	175	148	84
AY353491	1500	282	1.00E-75	199	154	77	AY353436	1488	152	2.00E-36	98	79	80
AY353491	1500	151	5.00E-36	98	78	79	AY353355	1521	281	3.00E-75	175	148	84
AY353287	1521	282	1.00E-75	275	164	59	AY353355	1521	155	3.00E-37	100	81	81
AY353287	1521	155	3.00E-37	100	81	81	AY353354	1503	281	3.00E-75	175	148	84
AY353286	1521	282	1.00E-75	275	164	59	AY353354	1503	157	9.00E-38	100	82	82
AY353286	1521	155	3.00E-37	100	81	81	AY353353	1503	281	3.00E-75	175	148	84
AY353285	1521	282	1.00E-75	275	164	59	AY353353	1503	157	9.00E-38	100	82	82
AY353285	1521	155	3.00E-37	100	81	81	AY353352	1503	281	3.00E-75	175	148	84
AJ430202	8179	282	2.00E-75	296	167	56	AY353352	1503	155	3.00E-37	100	81	81
AY353527	1521	282	2.00E-75	217	156	71	AY353330	1488	281	3.00E-75	175	148	84
AY353527	1521	154	5.00E-37	100	80	80	AY353330	1488	155	3.00E-37	100	81	81
AY353300	1521	282	2.00E-75	232	159	68	AY353329	1488	281	3.00E-75	175	148	84
AY353300	1521	155	3.00E-37	100	81	81	AY353329	1488	155	3.00E-37	100	81	81
AY353299	1521	282	2.00E-75	232	159	68	AY353328	1488	281	3.00E-75	175	148	84
AY353299	1521	155	3.00E-37	100	81	81	AY353328	1488	155	3.00E-37	100	81	81
AY353298	1521	282	2.00E-75	232	159	68	AY353327	1506	281	3.00E-75	175	148	84
AY353298	1521	155	3.00E-37	100	81	81	AY353327	1506	155	3.00E-37	100	81	81
AY353297	1521	282	2.00E-75	232	159	68	AY353326	1506	281	3.00E-75	175	148	84
AY353297	1521	155	3.00E-37	100	81	81	AY353326	1506	155	3.00E-37	100	81	81
AY353296	1521	282	2.00E-75	232	159	68	AY353325	1506	281	3.00E-75	175	148	84
AY353296	1521	155	3.00E-37	100	81	81	AY353325	1506	155	3.00E-37	100	81	81
AY353277	1521	282	2.00E-75	232	159	68	AY353324	1506	281	3.00E-75	175	148	84
AY353277	1521	155	3.00E-37	100	81	81	AY353324	1506	155	3.00E-37	100	81	81
AY353276	1521	282	2.00E-75	232	159	68	AY353323	1506	281	3.00E-75	175	148	84
AY353276	1521	155	3.00E-37	100	81	81	AY353323	1506	155	3.00E-37	100	81	81
AY353275	1521	282	2.00E-75	232	159	68	AY353320	1506	281	3.00E-75	175	148	84

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AY353320	1506	155	3.00E-37	100	81	81	AY649730	1482	152	2.00E-36	98	79	80
AY353319	1506	281	3.00E-75	175	148	84	AY649729	1482	280	5.00E-75	175	148	84
AY353319	1506	155	3.00E-37	100	81	81	AY649729	1482	152	2.00E-36	98	79	80
AY353318	1506	281	3.00E-75	175	148	84	AY649728	1482	280	5.00E-75	175	148	84
AY353318	1506	155	3.00E-37	100	81	81	AY649728	1482	152	2.00E-36	98	79	80
AY353317	1506	281	3.00E-75	175	148	84	AY649727	1482	280	5.00E-75	175	148	84
AY353317	1506	155	3.00E-37	100	81	81	AY649727	1482	152	2.00E-36	98	79	80
AY353316	1506	281	3.00E-75	175	148	84	AY649726	1482	280	5.00E-75	175	148	84
AY353316	1506	155	3.00E-37	100	81	81	AY649726	1482	152	2.00E-36	98	79	80
AY353315	1506	281	3.00E-75	175	148	84	AY649722	1488	280	5.00E-75	175	148	84
AY353315	1506	155	3.00E-37	100	81	81	AY649722	1488	156	1.00E-37	245	106	43
AY353314	1506	281	3.00E-75	175	148	84	AY649721	1488	280	5.00E-75	175	148	84
AY353314	1506	155	3.00E-37	100	81	81	AY649721	1488	152	2.00E-36	98	79	80
AY353311	1506	281	3.00E-75	175	148	84	AY649720	1488	280	5.00E-75	175	148	84
AY353311	1506	155	3.00E-37	100	81	81	AY649720	1488	152	2.00E-36	98	79	80
AY353309	1506	281	3.00E-75	175	148	84	AY649719	1488	280	5.00E-75	175	148	84
AY353309	1506	155	3.00E-37	100	81	81	AY649719	1488	152	2.00E-36	98	79	80
AY353308	1506	281	3.00E-75	175	148	84	AY649718	1488	280	5.00E-75	175	148	84
AY353308	1506	155	3.00E-37	100	81	81	AY649718	1488	152	2.00E-36	98	79	80
AY353306	1506	281	3.00E-75	175	148	84	AY649699	1506	280	5.00E-75	175	148	84
AY353306	1506	155	3.00E-37	100	81	81	AY649699	1506	154	5.00E-37	100	81	81
AY353305	1506	281	3.00E-75	175	148	84	AY649697	1488	280	5.00E-75	175	148	84
AY353305	1506	155	3.00E-37	100	81	81	AY649697	1488	156	1.00E-37	100	82	82
AY353295	1521	281	3.00E-75	175	148	84	AE008787	24186	280	5.00E-75	175	148	84
AY353295	1521	155	3.00E-37	100	81	81	AE008787	24186	152	2.00E-35	98	79	80
AY353294	1521	281	3.00E-75	175	148	84	X03394	1482	280	5.00E-75	175	148	84
AY353294	1521	155	3.00E-37	100	81	81	X03394	1482	154	8.00E-37	329	126	38
AY353293	1398	281	3.00E-75	175	148	84	X03393	1497	280	5.00E-75	175	148	84
AY353293	1398	155	3.00E-37	100	81	81	X03393	1497	157	5.00E-38	100	83	83
AY353289	1521	281	3.00E-75	175	148	84	AY657000	1474	280	5.00E-75	175	148	84
AY353289	1521	155	3.00E-37	100	81	81	AY657000	1474	138	1.00E-31	90	71	78
AY353284	1521	281	3.00E-75	175	148	84	X04505	1479	280	5.00E-75	175	148	84
AY353284	1521	155	3.00E-37	100	81	81	X04505	1479	152	2.00E-36	98	79	80
AY353282	1521	281	3.00E-75	175	148	84	AF338929	1506	280	5.00E-75	175	148	84
AY353282	1521	155	3.00E-37	100	81	81	AF335929	1506	154	8.00E-37	100	80	80
AY353280	1521	281	3.00E-75	175	148	84	AF420425	1460	280	5.00E-75	175	148	84
AY353280	1521	155	3.00E-37	100	81	81	AF420425	1460	129	3.00E-29	85	66	77
U17174	1521	281	3.00E-75	175	148	84	AY353526	1521	280	5.00E-75	175	148	84
U17174	1521	155	3.00E-37	100	81	81	AY353526	1521	155	3.00E-37	100	81	81
U17171	1521	281	3.00E-75	175	148	84	AY353519	1503	280	5.00E-75	175	148	84
U17171	1521	154	5.00E-37	100	81	81	AY353519	1503	152	2.00E-36	98	79	80
AY353462	1482	281	4.00E-75	229	160	69	AY353515	1488	280	5.00E-75	175	148	84
AY353462	1482	147	7.00E-35	98	77	78	AY353515	1488	156	1.00E-37	245	106	43
AY353461	1482	281	4.00E-75	229	160	69	AY353513	1491	280	5.00E-75	175	148	84
AY353461	1482	152	2.00E-36	98	79	80	AY353513	1491	155	3.00E-37	100	81	81
AY353460	1482	281	4.00E-75	229	160	69	AY353511	1491	280	5.00E-75	175	148	84
AY353460	1482	152	2.00E-36	98	79	80	AY353511	1491	153	1.00E-36	100	80	80
AY353339	1500	281	4.00E-75	217	157	72	AY353510	1491	280	5.00E-75	175	148	84
AY353339	1500	155	3.00E-37	100	81	81	AY353510	1491	152	2.00E-36	98	79	80
AY353338	1500	281	4.00E-75	217	157	72	AY353481	1488	280	5.00E-75	175	148	84
AY353338	1500	155	3.00E-37	100	81	81	AY353481	1488	154	5.00E-37	100	81	81
AY353337	1500	281	4.00E-75	217	157	72	AY353480	1488	280	5.00E-75	175	148	84
AY353337	1500	155	3.00E-37	100	81	81	AY353480	1488	152	2.00E-36	98	79	80
AY353336	1500	281	4.00E-75	217	157	72	AY353472	1500	280	5.00E-75	175	148	84
AY353336	1500	155	3.00E-37	100	81	81	AY353472	1500	154	6.00E-37	98	80	81
AY353335	1500	281	4.00E-75	217	157	72	AY353471	1500	280	5.00E-75	175	148	84
AY353335	1500	155	3.00E-37	100	81	81	AY353471	1500	154	6.00E-37	98	80	81
AY649740	1506	280	5.00E-75	175	148	84	AY353468	1500	155	3.00E-37	100	81	81
AY649740	1506	154	5.00E-37	100	81	81	AY353468	1500	280	5.00E-75	175	148	84
AY649739	1500	280	5.00E-75	175	148	84	AY353467	1500	155	3.00E-37	100	81	81
AY649739	1500	155	3.00E-37	100	81	81	AY353467	1500	155	3.00E-37	100	81	81
AY649734	1500	280	5.00E-75	175	148	84	AY353466	1521	280	5.00E-75	175	148	84
AY649734	1500	154	6.00E-37	98	80	81	AY353466	1521	159	2.00E-38	100	83	83
AY649733	1488	280	5.00E-75	175	148	84	AY353465	1521	280	5.00E-75	175	148	84
AY649733	1488	152	2.00E-36	98	79	80	AY353465	1521	158	3.00E-38	98	82	83
AY649731	1482	280	5.00E-75	175	148	84	AY353464	1476	280	5.00E-75	175	148	84
AY649731	1482	152	2.00E-36	98	79	80	AY353464	1476	152	2.00E-36	98	79	80
AY649730	1482	280	5.00E-75	175	148	84	AY353463	1476	280	5.00E-75	175	148	84

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#ident	%S
AY353463	1476	152	2.00E-36	98	79	80	AY353322	1506	155	3.00E-37	100	81	81
AY353459	1482	280	5.00E-75	175	148	84	AY353321	1506	280	5.00E-75	175	148	84
AY353459	1482	152	2.00E-36	98	79	80	AY353321	1506	155	3.00E-37	100	81	81
AY353458	1482	280	5.00E-75	175	148	84	AY353313	1506	280	5.00E-75	175	148	84
AY353458	1482	152	2.00E-36	98	79	80	AY353313	1506	155	3.00E-37	100	81	81
AY353442	1488	280	5.00E-75	175	148	84	AY353312	1506	280	5.00E-75	175	148	84
AY353442	1488	156	1.00E-37	245	106	43	AY353312	1506	155	3.00E-37	100	81	81
AY353441	1488	280	5.00E-75	175	148	84	AY353307	1506	280	5.00E-75	175	148	84
AY353441	1488	157	9.00E-38	245	106	43	AY353307	1506	155	3.00E-37	100	81	81
AY353432	1488	280	5.00E-75	175	148	84	AY353301	1488	280	5.00E-75	175	148	84
AY353432	1488	152	2.00E-36	98	79	80	AY353301	1488	155	3.00E-37	100	81	81
AY353431	1488	280	5.00E-75	175	148	84	AY353291	1521	280	5.00E-75	175	148	84
AY353431	1488	152	2.00E-36	98	79	80	AY353291	1521	154	5.00E-37	98	80	81
AY353430	1488	280	5.00E-75	175	148	84	AY353290	1521	280	5.00E-75	175	148	84
AY353430	1488	152	2.00E-36	98	79	80	AY353290	1521	154	5.00E-37	98	80	81
AY353429	1488	280	5.00E-75	175	148	84	AY353281	1521	280	5.00E-75	175	148	84
AY353429	1488	152	2.00E-36	98	79	80	AY353281	1521	154	5.00E-37	98	80	81
AY353428	1488	280	5.00E-75	175	148	84	AY353279	1521	280	5.00E-75	175	148	84
AY353428	1488	152	2.00E-36	98	79	80	AY353279	1521	154	5.00E-37	98	80	81
AY353427	1488	280	5.00E-75	175	148	84	AY353278	1521	280	5.00E-75	175	148	84
AY353427	1488	152	2.00E-36	98	79	80	AY353278	1521	154	5.00E-37	98	80	81
AY353426	1488	280	5.00E-75	175	148	84	AY353274	1521	280	5.00E-75	175	148	84
AY353426	1488	152	2.00E-36	98	79	80	AY353274	1521	154	5.00E-37	98	80	81
AY353425	1488	280	5.00E-75	175	148	84	AY353273	1521	280	5.00E-75	175	148	84
AY353425	1488	152	2.00E-36	98	79	80	AY353273	1521	154	5.00E-37	98	80	81
AY353424	1488	280	5.00E-75	175	148	84	AY353262	1488	280	5.00E-75	175	148	84
AY353424	1488	152	2.00E-36	98	79	80	AY353262	1488	155	3.00E-37	100	81	81
AY353423	1488	280	5.00E-75	175	148	84	AF332501	1515	280	5.00E-75	173	148	85
AY353423	1488	152	2.00E-36	98	79	80	AF332601	1515	149	2.00E-35	91	76	83
AY353422	1488	280	5.00E-75	175	148	84	D13689	1826	280	5.00E-75	175	148	84
AY353422	1488	152	2.00E-36	98	79	80	D13689	1826	152	2.00E-36	98	79	80
AY353421	1488	280	5.00E-75	175	148	84	AF159459	1602	280	7.00E-75	175	148	84
AY353421	1488	152	2.00E-36	98	79	80	AF159459	1602	154	5.00E-37	100	81	81
AY353420	1488	280	5.00E-75	175	148	84	AY353440	1488	280	9.00E-75	175	147	84
AY353420	1488	152	2.00E-36	98	79	80	AY353440	1488	152	2.00E-36	98	79	80
AY353419	1488	280	5.00E-75	175	148	84	AY353261	1485	280	9.00E-75	175	147	84
AY353419	1488	152	2.00E-36	98	79	80	AY353261	1485	153	1.00E-36	100	80	80
AY353418	1488	280	5.00E-75	175	148	84	AY353260	1485	280	9.00E-75	175	147	84
AY353418	1488	152	2.00E-36	98	79	80	AY353260	1485	152	2.00E-36	98	79	80
AY353417	1488	280	5.00E-75	175	148	84	AY374784	1472	194	9.00E-75	114	97	85
AY353417	1488	152	2.00E-36	98	79	80	AY374784	1472	163	1.00E-39	258	114	44
AY353369	1506	280	5.00E-75	175	148	84	AY374784	1472	113	9.00E-75	66	60	90
AY353369	1506	152	2.00E-36	98	79	80	D13690	3471	279	1.00E-74	175	147	84
AY353368	1506	280	5.00E-75	175	148	84	D13690	3471	155	3.00E-37	100	81	81
AY353368	1506	154	5.00E-37	100	81	81	AY649738	1491	279	2.00E-74	175	147	84
AY353367	1506	280	5.00E-75	175	148	84	AY649738	1491	153	1.00E-36	100	80	80
AY353367	1506	154	5.00E-37	100	81	81	AY249996	1368	279	2.00E-74	245	160	65
AY353366	1506	280	5.00E-75	175	148	84	AY249996	1368	142	2.00E-33	90	73	81
AY353366	1506	152	2.00E-36	98	79	80	AY353260	1503	279	2.00E-74	175	147	84
AY353365	1506	280	5.00E-75	175	148	84	AY353260	1503	153	1.00E-36	100	80	80
AY353365	1506	154	5.00E-37	100	81	81	AY353473	1500	279	2.00E-74	175	147	84
AY353364	1506	280	5.00E-75	175	148	84	AY353473	1500	152	2.00E-36	98	79	80
AY353364	1506	154	5.00E-37	100	81	81	AY353265	1521	279	2.00E-74	232	158	68
AY353357	1488	280	5.00E-75	175	148	84	AY353265	1521	155	3.00E-37	100	81	81
AY353357	1488	156	1.00E-37	100	82	82	AY353258	1494	279	2.00E-74	175	147	84
AY353356	1488	280	5.00E-75	175	148	84	AY353258	1494	155	2.00E-37	273	112	41
AY353356	1488	156	1.00E-37	100	82	82	AY353370	1506	278	2.00E-74	175	147	84
AY353351	1497	280	5.00E-75	175	148	84	AY353370	1506	152	2.00E-36	98	79	80
AY353351	1497	155	3.00E-37	100	81	81	AY353344	1488	278	2.00E-74	175	147	84
AY353343	1488	280	5.00E-75	175	148	84	AY353344	1488	155	3.00E-37	100	81	81
AY353343	1488	155	3.00E-37	100	81	81	AB128921	2358	278	3.00E-74	158	145	91
AY353342	1488	280	5.00E-75	175	148	84	AB128921	2358	136	1.00E-31	85	72	84
AY353342	1488	155	3.00E-37	100	81	81	AY249993	1383	278	4.00E-74	244	157	64
AY353342	1488	280	5.00E-75	175	148	84	AY249993	1383	142	2.00E-33	90	73	81
AY353341	1488	155	3.00E-37	100	81	81	AY353437	1488	278	4.00E-74	175	147	84
AY353341	1488	155	3.00E-37	100	81	81	AY353437	1488	152	2.00E-36	98	79	80
AY353340	1488	280	5.00E-75	175	148	84	M11332	1485	277	5.00E-74	175	146	83
AY353340	1488	155	3.00E-37	100	81	81	M11332	1485	153	1.00E-36	100	80	80
AY353322	1506	280	5.00E-75	175	148	84							

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AF128953	1668	276	1.00E-73	258	154	59	AF077601	831	243	1.00E-63	292	136	46
AF128953	1668	123	1.00E-27	103	64	62	AF198617	6830	242	2.00E-63	304	141	46
AF128950	1681	276	1.00E-73	258	154	59	AF198617	6830	238	4.00E-62	305	142	46
AF128950	1681	132	3.00E-30	107	68	63	AF105060	3765	236	1.00E-61	296	129	43
AF487406	1656	275	2.00E-73	222	151	68	AE015761	14255	236	2.00E-61	297	140	47
AF487406	1656	143	1.00E-33	335	118	35	AE015761	14255	233	1.00E-60	297	138	46
AY250009	1365	275	2.00E-73	238	159	66	AE015761	14255	55.1	5.00E-07	136	36	26
AY250009	1365	140	9.00E-33	90	72	80	AE015761	14255	34.3	0.89	89	26	29
AF517662	1348	275	3.00E-73	243	158	65	BA0000433544776	234	4.00E-61	307	134	43	
AF517662	1348	126	2.00E-28	85	65	76	BA0000433544776	142	2.00E-33	269	96	35	
AY250004	1380	274	4.00E-73	158	141	89	BA0000433544776	103	2.00E-21	179	62	34	
AY250004	1380	142	2.00E-33	90	73	81	BA0000433544776	80.5	1.00E-14	288	69	23	
AF517661	1361	274	4.00E-73	242	155	64	AB061233	849	233	1.00E-60	299	137	45
AF517661	1361	124	7.00E-26	84	64	76	AF077600	834	232	2.00E-60	296	137	46
AY250000	1506	273	7.00E-73	281	166	59	AF399739	6492	231	5.00E-60	300	138	46
AY250000	1506	144	8.00E-34	88	73	82	AF232939	1045	230	6.00E-60	348	157	45
AF128952	1695	273	1.00E-72	208	143	68	AF232941	1051	230	8.00E-60	350	154	44
AF128952	1695	117	1.00E-25	99	60	60	AY590686	874	229	1.00E-59	289	127	43
AF517665	1365	272	1.00E-72	158	140	88	AY590679	873	229	1.00E-59	289	127	43
AF517665	1365	130	9.00E-30	294	104	35	AY590678	873	229	1.00E-59	289	127	43
AY353514	1500	272	1.00E-72	197	149	75	AY590677	873	229	1.00E-59	289	127	43
AY353514	1500	152	2.00E-36	98	79	80	AY590676	873	229	1.00E-59	289	127	43
BX640427	348997	272	1.00E-72	392	180	45	AY590675	873	229	1.00E-59	289	127	43
BX640427	348997	54.3	8.00E-07	136	42	30	AY590674	864	229	1.00E-59	289	127	43
AF128948	1569	272	2.00E-72	251	156	62	AY590673	873	229	1.00E-59	289	127	43
AF128948	1569	128	5.00E-29	93	62	66	AY590672	873	229	1.00E-59	289	127	43
AF517664	1343	271	3.00E-72	236	157	66	AY590671	873	229	1.00E-59	289	127	43
AF517664	1343	119	2.00E-26	84	61	72	AY590670	873	229	1.00E-59	289	127	43
AF517663	1356	270	7.00E-72	156	139	89	AY590669	873	229	1.00E-59	289	127	43
AF517663	1356	124	7.00E-28	84	64	76	AE0171803814139	228	3.00E-59	298	133	44	
AF128952	1665	268	3.00E-71	253	151	59	AE0171803814139	90.1	1.00E-17	293	76	25	
AF128952	1665	132	3.00E-30	107	68	63	AB128920	2876	228	3.00E-59	127	121	95
AF128951	1665	268	3.00E-71	253	151	59	AB128920	2876	167	7.00E-41	313	124	39
AF128951	1665	132	3.00E-30	107	68	63	AB128920	2876	85.9	3.00E-16	51	40	78
BX640413	349028	268	4.00E-71	391	182	46	AB061232	849	227	5.00E-59	299	131	43
BX640444	349008	266	1.00E-70	391	183	46	AB105910	15284	226	1.00E-58	299	131	43
BX640444	349008	54.3	8.00E-07	136	42	30	AB061230	19187	226	1.00E-58	299	131	43
L13034	1572	266	1.00E-70	391	183	46	AB061230	19187	59.3	3.00E-08	130	38	29
AF128954	1680	265	2.00E-70	204	140	68	AE016862	310286	223	8.00E-58	297	129	43
AF128954	1680	114	9.00E-25	97	58	59	AF095238	846	222	2.00E-57	291	129	44
AF128947	1649	265	2.00E-70	254	149	58	AB061231	849	222	2.00E-57	297	129	43
AF128947	1649	117	6.00E-26	100	62	62	AE013021	10029	220	7.00E-57	298	134	44
AF128955	1524	265	3.00E-70	247	153	61	BA000021	697724	219	1.00E-56	181	109	60
AF128955	1524	112	3.00E-24	82	53	64	BA000021	697724	120	9.00E-27	345	109	31
AY249136	1398	263	7.00E-70	160	137	85	AF077341	874	219	1.00E-56	292	124	42
AY249136	1398	138	4.00E-32	161	84	52	AF095237	873	219	2.00E-56	292	123	42
AF128956	1551	262	2.00E-69	245	152	62	AB080999	348	218	3.00E-56	115	115	100
AF128956	1551	125	1.00E-28	93	62	66	AB033501	1210	217	6.00E-56	330	134	40
AF128949	1563	259	2.00E-68	246	151	61	CP000024222334	215	3.00E-55	313	135	43	
AF128949	1563	121	6.00E-27	90	58	64	CP000024222334	33.9	1.2	87	22	25	
AF425736	1351	254	3.00E-67	160	133	83	AE017334222645	215	3.00E-55	313	135	43	
AF425736	1351	92.4	3.00E-18	64	46	71	AE017334222645	33.9	1.2	87	22	25	
AF128957	1668	254	3.00E-67	248	143	57	BA0000444202352	214	6.00E-55	294	124	42	
AF128957	1668	129	2.00E-29	106	67	63	BA0000444202352	130	1.00E-29	135	70	51	
AY534751	1404	253	7.00E-67	158	135	85	BA0000444202352	95.9	2.00E-19	169	57	33	
AY534751	1404	123	1.00E-27	117	70	59	D10063	1064	214	6.00E-55	294	124	42
AY250005	1383	251	4.00E-66	177	133	75	AF232940	1045	213	1.00E-54	273	137	50
AY250008	1383	118	5.00E-26	280	93	33	AF065259	1631	211	3.00E-54	300	125	41
AY534750	1380	250	8.00E-66	158	133	84	AF095236	873	211	3.00E-54	300	125	41
AY534750	1380	123	1.00E-27	189	82	43	AF011370	1300	211	5.00E-54	381	141	37
AF002709	3434	250	8.00E-66	303	140	46	AE015942	300171	209	1.00E-53	296	125	42
AF002709	3434	239	2.00E-62	304	139	45	AE015942	300171	68.2	5.00E-11	284	68	23
AY250007	1197	249	2.00E-65	158	127	80	BA0000283630528	209	1.00E-53	338	131	38	
AY250007	1197	107	6.00E-23	304	93	30	BA0000283630528	91.3	6.00E-18	293	75	25	
AE016923	303642	248	4.00E-65	297	137	46	AF080260	1149	209	2.00E-53	380	141	37
AE016923	303642	246	1.00E-64	297	137	46	AF080259	1149	209	2.00E-53	380	141	37
AF420426	1364	248	4.00E-65	157	130	82	Z99122	200690	208	3.00E-53	306	130	42
AF420426	1364	114	5.00E-25	77	58	75	U56901	20320	208	3.00E-53	306	130	42

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
M26948	2360	208	3.00E-53	306	130	42	AF084813	1952	189	1.00E-47	385	136	35
M26947	1474	208	3.00E-53	306	130	42	AF084812	1952	189	1.00E-47	385	136	35
AB058936	884	207	4.00E-53	294	123	41	AFD19213	7256	189	1.00E-47	375	130	34
BA0000321877212	207	6.00E-53	301	119	39	AF030239	1167	189	1.00E-47	385	136	35	
BA0000321877212	36.6	0.18	116	29	25	U82287	1199	189	1.00E-47	385	136	35	
AF124349	42050	207	6.00E-53	299	120	40	U82286	1199	189	1.00E-47	385	136	35
AE007720	7354	207	8.00E-53	295	122	41	AE017003	347456	189	2.00E-47	296	114	38
AF064086	1020	207	8.00E-53	295	122	41	AE017003	347456	180	1.00E-44	298	115	38
AE0086922056416	206	1.00E-52	296	119	40	AE017003	347456	165	3.00E-40	271	98	36	
U52957	9544	206	2.00E-52	301	119	39	AE017003	347456	140	9.00E-33	291	95	32
L06176	3724	206	2.00E-52	301	119	39	AE017003	347456	56.6	2.00E-07	243	57	23
BA0000313288558	201	3.00E-51	376	131	34	AF111790	1161	189	2.00E-47	384	136	35	
BA0000313288558	173	9.00E-43	384	125	32	AF098793	1167	189	2.00E-47	385	134	34	
BA0000313288558	167	9.00E-41	377	125	33	AF078151	1161	189	2.00E-47	384	136	35	
BA0000313288558	167	9.00E-41	377	126	33	AP0068403566135	189	2.00E-47	296	114	38		
BA0000313288558	167	9.00E-41	377	126	33	AF078152	1181	188	3.00E-47	384	136	35	
BA0000313288558	109	2.00E-23	148	60	40	AJ748315	3011	188	3.00E-47	301	117	38	
BA0000313288558	74.3	8.00E-13	217	61	28	AE017029	290525	188	4.00E-47	292	112	38	
AB058937	820	199	1.00E-50	287	118	41	AE0173345227419	188	4.00E-47	292	112	38	
AJ748317	3009	199	2.00E-50	295	124	42	AE0172255228663	188	4.00E-47	292	112	38	
BX842647	346357	199	2.00E-50	295	124	42	Y18889	5040	187	5.00E-47	300	111	37
BX842647	346357	196	2.00E-49	292	120	41	Y18889	5040	174	7.00E-43	304	103	33
BX842647	346357	188	3.00E-47	301	117	38	AF078154	1161	187	5.00E-47	384	135	35
BX842647	346357	186	2.00E-46	292	115	39	AF007121	4400	187	5.00E-47	375	129	34
AE011832	13074	196	2.00E-49	212	113	53	AF007121	4400	175	2.00E-43	377	129	34
AE011832	13074	103	9.00E-22	307	87	28	AE017251	301045	187	6.00E-47	300	113	37
AE011832	13074	62.4	3.00E-09	134	40	29	AE017251	301045	172	3.00E-42	304	106	34
AB039913	921	195	3.00E-49	302	118	39	AB110834	1155	187	8.00E-47	382	131	34
AF081500	1152	194	4.00E-49	380	136	35	AJ851165	2169	186	2.00E-46	292	115	39
AF078155	1146	194	5.00E-49	379	136	35	X67138	5206	185	3.00E-46	292	107	36
AF078153	1146	194	5.00E-49	379	136	35	X67138	5206	182	3.00E-45	292	107	36
AJ748319	3196	194	5.00E-49	291	119	40	X67138	5206	180	1.00E-44	296	107	36
AJ748319	3196	90.1	1.00E-17	101	49	48	AB039909	750	185	3.00E-46	269	114	42
AE012298	12693	193	9.00E-49	241	119	49	U52198	5354	184	5.00E-46	375	128	34
AE012298	12693	104	7.00E-22	319	93	29	U52198	5354	155	3.00E-37	182	80	43
AE012298	12693	58.2	6.00E-08	134	38	28	U52198	5354	150	9.00E-36	376	119	31
AY514454	9258	193	9.00E-49	379	140	36	AB040140	1479	184	5.00E-46	235	116	49
AY514454	9258	174	4.00E-43	370	126	34	AB040140	1479	114	9.00E-25	251	80	31
AY514454	9258	174	4.00E-43	377	126	33	AB039906	756	184	5.00E-46	270	111	41
AY514454	9258	153	1.00E-36	218	89	40	CR378655347213	184	5.00E-46	382	130	34	
AY514454	9258	57	1.00E-07	60	27	45	CR378655347213	145	3.00E-34	213	82	38	
AB039905	921	193	9.00E-49	302	117	38	CR378655347213	90.5	1.00E-17	203	63	31	
AB039938	817	192	1.00E-48	287	119	41	AE001250	14793	183	1.00E-45	298	114	38
CR5228703523383	192	2.00E-48	252	120	47	AY331139	688	182	2.00E-45	251	102	40	
CR5228703523383	177	8.00E-44	231	108	46	AB110832	1155	182	2.00E-45	381	132	34	
CR5228703523383	115	3.00E-25	227	81	35	AE007665	12205	182	2.00E-45	292	115	39	
CR5228703523383	107	1.00E-22	254	79	31	AE017314	300029	182	2.00E-45	311	119	38	
CR5228703523383	51.6	5.00E-06	130	40	30	AE004287	14585	182	3.00E-45	375	133	35	
CR5228703523383	34.3	0.89	148	33	22	AE004287	14585	176	1.00E-43	377	134	35	
AB039911	927	191	3.00E-48	304	119	39	AE004287	14585	169	1.00E-41	379	127	33
AB039907	927	191	4.00E-46	304	119	39	AE069392	39101	182	3.00E-45	376	124	32
AE011409	10859	190	7.00E-48	302	111	36	AE069392	39101	167	9.00E-41	377	125	33
AE011409	10859	176	2.00E-43	300	107	35	AE069392	39101	167	9.00E-41	377	126	33
AE017293	301124	190	7.00E-48	302	111	36	AE007122	4700	182	3.00E-45	375	133	35
AE017293	301124	176	1.00E-43	300	107	35	AE007122	4700	176	1.00E-43	377	134	35
AE017293	301124	33.9	1.2	59	18	30	AE007122	4700	171	6.00E-42	379	127	33
AB039910	927	190	1.00E-47	304	119	39	X63965	2072	181	3.00E-45	301	112	37
M20983	1031	190	1.00E-47	304	121	39	X63965	2072	181	6.00E-45	304	107	35
AE004290	10581	189	1.00E-47	375	130	34	AB040139	1479	181	3.00E-45	236	111	47
AE004290	10581	175	2.00E-43	377	128	34	AB040139	1479	115	3.00E-25	284	83	29
CP0000103510148	189	1.00E-47	385	136	35	AY331140	686	181	6.00E-45	251	105	41	
CP0000103510148	34.3	0.89	76	24	31	AE001257	13979	181	6.00E-45	301	111	36	
BX5719654074542	189	1.00E-47	385	136	35	AE001257	13979	181	6.00E-45	304	107	35	
BX5719654074542	40	0.016	335	74	22	AY331141	686	180	8.00E-45	251	105	41	
BX5719654074542	34.3	0.89	76	24	31	M94015	1198	180	8.00E-45	304	113	37	
U73848	2046	189	1.00E-47	385	136	35	AB110833	1155	180	1.00E-44	381	131	34
AF084815	1951	189	1.00E-47	385	136	35	AB110831	1155	179	1.00E-44	381	131	34
AF084814	1951	189	1.00E-47	385	136	35	CR6283363503610	179	1.00E-44	309	124	40	

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	
CR6283363503610	110	7.00E-24	200	77	38		AF320637	1685	174	5.00E-43	300	107	35	
L47122	2101	1.00E-44	379	129	34		L42885	1011	174	5.00E-43	342	113	33	
L34686	2350	1.00E-44	304	116	38		X69614	1011	174	7.00E-43	342	112	32	
AE0173555237682	179	2.00E-44	365	125	34		X69810	1011	174	7.00E-43	342	113	33	
AE0173555237682	142	3.00E-33	292	97	33		AB022136	974	174	7.00E-43	329	114	34	
AY166716	1608	2.00E-44	335	120	35		AE016803	300045	173	9.00E-43	375	121	32	
AY028400	8125	2.00E-44	296	109	36		AE016803	300045	169	1.00E-41	376	122	32	
AF348135	3834	2.00E-44	301	109	36		AE016803	300045	151	5.00E-36	376	119	31	
AF348135	3834	2.00E-42	301	109	36		AY560550	864	173	9.00E-43	296	107	36	
AE0173543397754	178	3.00E-44	309	125	40		AY560549	864	173	9.00E-43	296	107	36	
AE0173543397754	109	2.00E-23	198	78	39		X63413	1011	173	9.00E-43	342	113	33	
AJ496278	1490	3.00E-44	309	123	39		X65624	1498	173	9.00E-43	296	107	36	
AJ496278	1490	2.00E-23	198	78	39		X16833	1435	173	9.00E-43	342	112	32	
AB022132	1011	3.00E-44	342	116	33		X15661	1011	173	9.00E-43	342	112	32	
AE011374	11076	6.00E-44	300	107	35		BA0000373354505	173	9.00E-43	375	121	32		
AE011374	11076	3.00E-39	301	100	33		BA0000373354505	153	1.00E-36	376	118	31		
AE017249	301384	6.00E-44	304	109	35		BA0000373354505	151	5.00E-36	376	119	31		
AE017294	300207	6.00E-44	300	107	35		BA0000373354505	149	2.00E-35	178	80	44		
AE017294	300207	4.00E-39	301	99	32		BA0000373354505	148	4.00E-35	376	111	29		
gi94646	107	2.00E-22	181	65	35		BA0000373354505	121	4.00E-27	374	108	28		
AB022136	1011	6.00E-44	342	115	33		BA0000373354505	102	3.00E-21	301	92	30		
AE016918	302178	8.00E-44	299	104	34		AL591976	250050	173	9.00E-43	296	107	36	
AE016918	302178	0.4	266	61	22		AL596166	260050	173	9.00E-43	296	107	36	
X69609	1011	8.00E-44	342	114	33		X69612	1011	173	9.00E-43	342	113	33	
AE017317	302040	8.00E-44	310	113	36		X69613	1008	173	9.00E-43	342	113	33	
AJ496382	1548	1.00E-43	309	122	39		AE017324	250242	173	9.00E-43	296	107	36	
AJ496382	1548	1.00E-23	251	88	35		AY275679	1223	173	1.00E-42	314	118	37	
AJ496283	1458	1.00E-43	309	122	39		AY275679	1223	122	3.00E-27	182	73	40	
AJ496283	1468	1.00E-23	251	88	35		AB022137	950	173	1.00E-42	323	113	34	
AJ496281	1472	1.00E-43	309	122	39		AE0168222584158	172	2.00E-42	299	105	35		
AJ496281	1472	95.9	2.00E-19	84	46	54		AE0168222584158	99.8	2.00E-20	308	85	27	
AJ496276	1493	1.00E-43	309	122	39		AJ496277	1486	172	2.00E-42	166	92	55	
AJ496276	1493	92.8	2.00E-18	138	61	44		AJ496277	1486	109	2.00E-23	198	78	39
X83232	1755	1.00E-43	309	122	39		AJ743318	3949	172	2.00E-42	295	109	36	
X83232	1755	2.00E-23	251	87	34		BX842654	344249	172	2.00E-42	295	109	36	
AJ496282	1482	1.00E-43	309	122	39		CR5553064296230	171	4.00E-42	151	89	58		
AJ496282	1482	1.00E-23	251	88	35		AB039912	843	171	4.00E-42	280	110	39	
CR6283373345687	176	1.00E-43	309	122	39		AY445112	990	171	5.00E-42	314	117	37	
CR6283373345687	110	1.00E-23	251	88	35		L81147	1272	171	5.00E-42	314	117	37	
ABC22133	1011	1.00E-43	342	115	33		D82864	973	171	5.00E-42	325	109	33	
AJ748316	3730	1.00E-43	295	110	37		AB014878	963	171	5.00E-42	323	111	34	
BX842655	349965	1.00E-43	295	110	37		AY660548	854	171	6.00E-42	298	106	35	
CP000013	904246	2.00E-43	342	113	33		AE007717	14157	171	6.00E-42	290	104	35	
CP000013	904246	0.52	77	18	23		AY275678	1223	171	6.00E-42	314	117	37	
AE01126	11037	2.00E-43	342	113	33		AY275678	1223	122	3.00E-27	182	73	40	
X69597	1011	2.00E-43	342	114	33		AY275677	1222	171	6.00E-42	314	117	37	
X69611	1008	2.00E-43	342	113	33		AY275677	1222	122	3.00E-27	182	73	40	
X69607	1008	2.00E-43	342	114	33		AB014677	973	171	6.00E-42	323	111	34	
AB039908	843	2.00E-43	285	113	39		AE015941	299511	170	6.00E-42	294	105	35	
X84699	2178	2.00E-43	228	105	46		AE015941	299511	142	2.00E-33	270	80	29	
X84699	2178	8.00E-23	214	79	36		X75200	1117	170	8.00E-42	342	111	32	
X53940	1005	2.00E-43	340	116	34		D82853	987	170	8.00E-42	335	111	33	
X75202	1123	2.00E-43	342	114	33		D82852	987	170	8.00E-42	335	111	33	
X15660	1011	2.00E-43	342	113	33		AE017316	300704	169	1.00E-41	295	101	34	
X56334	1426	2.00E-43	342	113	33		D82856	987	169	1.00E-41	335	110	32	
AL646078	203050	2.00E-43	297	109	36		AJ496279	1495	169	2.00E-41	314	122	38	
AF241832	1300	2.00E-43	296	107	36		AJ496279	1495	109	2.00E-23	198	78	39	
L42881	1011	1.00E-43	342	113	33		X75204	1121	169	2.00E-41	334	112	33	
L42876	1011	2.00E-43	342	113	33		L81146	1272	169	2.00E-41	300	114	38	
D43777	1398	2.00E-43	337	115	34		U54775	2745	168	3.00E-41	242	104	42	
M86838	1736	2.00E-43	340	116	34		U54775	2745	113	1.00E-24	335	100	29	
X69608	1006	3.00E-43	342	113	33		AY380808	1011	168	4.00E-41	342	110	32	
AF283285	1371	3.00E-43	300	111	37		D82855	987	168	4.00E-41	335	108	32	
AJ496275	1487	4.00E-43	310	122	39		AE016920	305584	167	5.00E-41	369	122	33	
AJ496275	1487	99.8	2.00E-20	191	72	37		AE016920	305584	58.9	3.00E-08	299	62	20
X75201	1112	174	4.00E-43	333	113	33		D82854	987	167	5.00E-41	335	108	32
X69598	1011	174	4.00E-43	342	113	33		D82848	987	167	5.00E-41	335	109	32
X75203	1087	174	5.00E-43	342	113	33		D82847	987	167	5.00E-41	335	109	32

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AB022134	987	167	5.00E-41	335	109	32	BA0000409105828	36.2	0.23	109	29	26	
AE004540	15833	167	7.00E-41	242	103	42	BA0000409105828	35.4	0.4	121	28	23	
AE004540	15833	113	2.00E-24	335	100	29	BA0000409105828	34.7	0.68	109	29	26	
AF034764	1170	167	7.00E-41	271	105	38	BA0000409105828	34.3	0.69	109	29	26	
AY275576	1229	167	7.00E-41	271	105	38	BA0000409105828	33.9	1.2	109	29	26	
AY275575	1228	167	7.00E-41	271	105	38	BA0000409105828	33.1	2	62	19	30	
AY275574	1243	167	7.00E-41	271	105	38	AF014114	852	160	1.00E-38	301	98	32
D82862	981	167	7.00E-41	333	109	32	AF034766	1458	160	1.00E-38	232	99	42
D82858	987	167	7.00E-41	335	108	32	AF034766	1458	100	8.00E-21	346	89	25
D82857	987	167	7.00E-41	335	108	32	AJ277361	822	159	2.00E-38	273	94	34
AF034768	1434	167	9.00E-41	225	101	44	D86618	984	159	2.00E-38	327	106	32
AF034768	1434	111	4.00E-24	278	83	29	D89073	1560	159	2.00E-38	256	101	39
D63372	987	167	9.00E-41	335	107	31	D89073	1560	89.7	2.00E-17	173	60	34
D63366	987	167	9.00E-41	335	108	32	D82859	984	159	2.00E-38	327	106	32
D63365	987	167	9.00E-41	335	108	32	AB058931	1313	159	2.00E-38	256	101	39
D63364	987	167	9.00E-41	335	107	31	AB058931	1313	89.7	2.00E-17	173	60	34
D82851	987	167	9.00E-41	335	107	31	AF034767	2007	159	2.00E-38	260	105	40
D82850	987	167	9.00E-41	335	107	31	AF034767	2007	108	3.00E-23	273	74	27
D82849	987	167	9.00E-41	335	107	31	AY424358	705	158	3.00E-38	217	92	42
AB022139	976	167	9.00E-41	323	109	33	AJ277358	822	158	3.00E-38	273	95	34
D63374	987	166	1.00E-40	335	107	31	AF084055	852	158	3.00E-38	301	98	32
D63373	987	166	1.00E-40	335	107	31	AY744156	858	158	4.00E-38	304	106	34
D63371	987	166	1.00E-40	335	107	31	AB022135	934	158	4.00E-38	313	103	32
D63370	987	166	1.00E-40	335	107	31	AB058932	1313	155	1.00E-37	278	102	36
D63369	987	166	1.00E-40	335	107	31	AB058932	1313	85.1	4.00E-16	108	45	41
D63368	987	166	1.00E-40	335	107	31	AJ277360	822	155	2.00E-37	273	93	34
D63367	987	166	1.00E-40	335	107	31	AF011371	1515	155	2.00E-37	246	97	39
D63363	987	166	1.00E-40	335	107	31	AF011371	1515	92	4.00E-18	87	46	52
D82846	987	166	1.00E-40	335	107	31	AB058934	1150	155	2.00E-37	374	115	30
D82851	981	166	1.00E-40	333	110	33	AJ277359	813	152	2.00E-36	268	91	33
M57501	1596	166	2.00E-40	271	105	38	BX572595	349250	152	2.00E-36	295	92	31
M57501	1596	107	1.00E-22	197	73	37	BX572595	349250	35	0.52	261	58	22
AF307102	981	165	3.00E-40	333	108	32	AE016797	301380	151	4.00E-36	376	120	31
AF307101	981	165	3.00E-40	333	108	32	AE016797	301380	149	2.00E-35	178	80	44
U12817	19811	165	3.00E-40	377	125	33	AE016797	301380	121	4.00E-27	374	108	28
U12817	19811	147	6.00E-35	188	78	41	AE016797	301380	102	3.00E-21	301	92	30
U12817	19811	107	6.00E-23	160	61	38	AB058933	1217	151	4.00E-36	169	82	48
U12817	19811	75.9	3.00E-13	217	59	27	AY192720	724	151	5.00E-36	263	96	36
U28496	1085	165	3.00E-40	334	111	33	AB058935	1156	151	5.00E-36	169	82	48
D82863	981	165	3.00E-40	333	108	32	AB058935	1156	79.7	2.00E-14	291	83	28
AE016790	300242	164	6.00E-40	252	103	40	CR378663	348044	151	5.00E-36	251	106	42
AE016790	300242	96.3	2.00E-19	129	53	41	CR378663	348044	89.4	2.00E-17	149	55	36
AE016790	300242	56.2	2.00E-07	130	41	31	U28499	1093	150	9.00E-36	335	102	30
L15366	2531	164	6.00E-40	252	103	40	AF011372	1620	150	1.00E-35	266	100	37
L15366	2531	95.9	2.00E-19	92	46	50	AF011372	1620	51.2	7.00E-06	147	43	29
AB017479	952	164	7.00E-40	323	108	33	U52199	2471	149	1.00E-35	376	116	30
AF064056	852	163	1.00E-39	301	100	33	AY192721	724	149	2.00E-35	263	95	36
AJ537492	1203	162	3.00E-39	103	84	81	AJ277355	661	149	2.00E-35	238	90	37
AJ537491	1202	162	3.00E-39	103	84	81	D90832	19562	149	2.00E-35	151	80	52
AJ537490	1203	162	3.00E-39	103	84	81	AB018734	603	149	2.00E-35	214	90	42
AJ537489	1204	162	3.00E-39	103	84	81	AE0173402839318	148	3.00E-35	170	82	48	
AJ537488	1203	162	3.00E-39	103	84	81	AE0173402839318	99.4	2.00E-20	267	77	28	
AJ537487	1202	162	3.00E-39	103	84	81	AE0173402839318	44.7	7.00E-04	130	35	26	
AJ537486	1201	162	3.00E-39	103	84	81	AJ277351	667	147	7.00E-35	237	85	35
AJ537485	1204	162	3.00E-39	103	84	81	AY192718	724	147	9.00E-35	267	96	35
X63513	1035	161	4.00E-39	307	109	35	AE017269	294300	145	3.00E-34	292	96	32
AF064057	852	161	5.00E-39	301	99	32	AE017269	294300	126	1.00E-28	221	77	34
M81344	1207	161	5.00E-39	301	99	32	AE017269	294300	75.1	5.00E-13	84	41	48
AF034765	1179	160	6.00E-39	300	108	36	AY192722	724	145	4.00E-34	263	91	34
BA0000409105828	160	8.00E-39	295	95	32	AJ277350	667	145	4.00E-34	237	84	35	
BA0000409105828	85.1	4.00E-16	311	73	23	U28498	1089	145	4.00E-34	330	102	30	
BA0000409105828	80.9	8.00E-16	317	74	23	AB018732	606	145	4.00E-34	217	91	41	
BA0000409105828	58.5	4.00E-08	273	72	26	AY192719	724	144	5.00E-34	264	93	35	
BA0000409105828	57.4	1.00E-07	255	72	28	AJ242662	1131	144	5.00E-34	377	112	29	
BA0000409105828	55.1	5.00E-07	250	63	25	AJ277356	660	144	6.00E-34	238	89	37	
BA0000409105828	53.9	1.00E-06	185	47	25	AY331135	1880	143	1.00E-33	263	99	37	
BA0000409105828	38.5	0.047	233	50	21	AY331135	1880	70.9	9.00E-12	68	35	51	
BA0000409105828	37.7	0.08	134	39	29	D85070	876	143	1.00E-33	291	94	32	

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
D85069	876	143	1.00E-33	291	94	32	AB027159	746	141	5.00E-33	267	87	32
AY192716	724	143	1.00E-33	264	92	34	AB027158	746	141	5.00E-33	267	87	32
AY192717	724	142	2.00E-33	264	89	33	AB027157	746	141	5.00E-33	267	87	32
D83764	876	142	2.00E-33	291	94	32	AY192726	724	140	7.00E-33	267	92	34
D83763	876	142	2.00E-33	291	94	32	AY192725	724	140	7.00E-33	267	92	34
D83762	876	142	2.00E-33	291	94	32	CP0000015300915	140	7.00E-33	292	95	32	
D88295	876	142	2.00E-33	291	93	31	CP0000015300915	137	6.00E-32	289	96	33	
AJ277352	667	142	2.00E-33	237	86	36	CP0000015300915	80.9	8.00E-15	111	46	41	
AF515472	674	142	2.00E-33	125	80	64	CP0000015300915	57.8	8.00E-08	242	57	23	
AF515472	674	57.8	8.00E-08	58	33	56	Y14687	2026	140	7.00E-33	166	74	44
D85071	876	142	3.00E-33	291	92	31	Y14687	2026	97.1	1.00E-19	176	63	35
D88291	866	142	3.00E-33	291	92	31	AF274346	699	140	7.00E-33	166	74	44
AB001704	876	142	3.00E-33	291	92	31	D85073	876	140	7.00E-33	291	92	31
AB167766	746	141	4.00E-33	267	88	32	AB174780	746	140	9.00E-33	267	87	32
X98463	998	141	4.00E-33	332	107	32	AB174779	746	140	9.00E-33	267	87	32
D85076	875	141	4.00E-33	291	92	31	AB174778	746	140	9.00E-33	267	87	32
D85075	876	141	4.00E-33	291	92	31	AB174777	746	140	9.00E-33	267	87	32
D85074	876	141	4.00E-33	291	92	31	AB174776	746	140	9.00E-33	267	87	32
D85072	876	141	4.00E-33	291	92	31	AB174775	746	140	9.00E-33	267	87	32
D88293	876	141	4.00E-33	291	92	31	AB174774	746	140	9.00E-33	267	87	32
AB001718	876	141	4.00E-33	291	92	31	AB031514	746	140	9.00E-33	267	87	32
AB001717	876	141	4.00E-33	291	92	31	AB027181	746	140	9.00E-33	267	87	32
AB001716	876	141	4.00E-33	291	92	31	AB027176	746	140	9.00E-33	267	87	32
AB001715	876	141	4.00E-33	291	92	31	AB027173	746	140	9.00E-33	267	87	32
AB001714	876	141	4.00E-33	291	92	31	AB027169	746	140	9.00E-33	267	87	32
AB001713	876	141	4.00E-33	291	92	31	AB027163	746	140	9.00E-33	267	87	32
AB001712	876	141	4.00E-33	291	92	31	AY192723	724	140	1.00E-32	267	92	34
AB001711	876	141	4.00E-33	291	92	31	AE001746	18364	140	1.00E-32	392	119	30
AB001710	876	141	4.00E-33	291	92	31	AF336830	772	140	1.00E-32	273	87	31
AB001709	876	141	4.00E-33	291	92	31	AB030272	746	140	1.00E-32	267	87	32
AB001708	876	141	4.00E-33	291	92	31	AB018737	600	139	2.00E-32	218	89	40
AB001707	876	141	4.00E-33	291	92	31	AB018736	600	139	2.00E-32	218	89	40
AB001706	876	141	4.00E-33	291	92	31	AY331136	1863	139	3.00E-32	258	96	37
AB001705	876	141	4.00E-33	291	92	31	AY331136	1863	63.2	2.00E-09	64	31	48
AB030271	746	141	4.00E-33	267	87	32	AY192727	722	139	3.00E-32	266	91	34
AB030270	746	141	4.00E-33	267	88	32	AJ277362	703	139	3.00E-32	248	85	34
AB027180	746	141	4.00E-33	267	88	32	AB058939	1685	138	4.00E-32	136	71	52
AB027175	746	141	4.00E-33	267	88	32	AB058939	1685	99.8	2.00E-20	291	82	28
AB027166	746	141	4.00E-33	267	87	32	AE007672	10661	137	7.00E-32	292	89	30
AB027161	746	141	4.00E-33	267	88	32	AY192724	723	135	2.00E-31	266	90	33
AB091716	746	141	5.00E-33	267	87	32	AY331137	1411	134	5.00E-31	233	86	36
AB031516	746	141	5.00E-33	267	87	32	AY331137	1411	65.1	5.00E-10	264	68	25
AB031515	746	141	5.00E-33	267	87	32	AF017113	47739	134	6.00E-31	206	81	39
AB031513	746	141	5.00E-33	267	87	32	Z99121	194692	134	6.00E-31	206	81	39
AB031512	746	141	5.00E-33	267	87	32	L15367	1666	130	7.00E-31	132	69	52
AB031511	746	141	5.00E-33	267	87	32	L15367	1666	97.4	9.00E-20	280	77	27
AB031510	746	141	5.00E-33	267	87	32	L15367	1666	29.6	7.00E-31	24	13	54
AB031509	746	141	5.00E-33	267	87	32	AB030273	746	134	8.00E-31	268	84	31
AB031508	746	141	5.00E-33	267	87	32	AJ277353	703	132	3.00E-30	248	83	33
AB018733	624	141	5.00E-33	214	87	40	AY551006	1380	132	3.00E-30	139	70	50
AB027188	746	141	5.00E-33	267	87	32	AY551006	1380	87.4	9.00E-17	159	57	35
AB027185	746	141	5.00E-33	267	87	32	AF026811	3031	131	4.00E-30	265	93	35
AB027184	746	141	5.00E-33	267	87	32	AF026811	3031	98.2	5.00E-20	153	53	41
AB027183	746	141	5.00E-33	267	87	32	AJ277363	669	131	5.00E-30	241	78	32
AB027182	746	141	5.00E-33	267	87	32	AB018735	600	130	9.00E-30	216	85	39
AB027179	746	141	5.00E-33	267	87	32	AY129557	9117	129	2.00E-29	251	82	32
AB027178	746	141	5.00E-33	267	87	32	AY129557	9117	49.3	3.00E-05	87	34	39
AB027177	746	141	5.00E-33	267	87	32	AJ277364	669	129	3.00E-29	241	79	32
AB027174	746	141	5.00E-33	267	87	32	AY551005	1482	129	3.00E-29	140	68	48
AB027172	746	141	5.00E-33	267	87	32	AY551005	1482	94	9.00E-19	195	67	34
AB027171	746	141	5.00E-33	267	87	32	AF026812	1863	128	5.00E-29	150	68	45
AB027170	746	141	5.00E-33	267	87	32	AF026812	1863	86.7	2.00E-16	158	58	36
AB027168	746	141	5.00E-33	267	87	32	AF312378	687	126	1.00E-28	248	79	31
AB027167	746	141	5.00E-33	267	87	32	AY331138	1293	124	7.00E-28	165	67	40
AB027165	746	141	5.00E-33	267	87	32	AY331138	1293	60.1	2.00E-08	92	33	35
AB027164	746	141	5.00E-33	267	87	32	AJ31736	2833	124	9.00E-28	81	63	77
AB027162	746	141	5.00E-33	267	87	32	AJ277357	670	122	3.00E-27	240	77	32
AB027160	746	141	5.00E-33	267	87	32	AY278534	577	122	3.00E-27	182	73	40

Fig. 25A-C-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AY278531	566	122	3.00E-27	132	73	40	U76543	1023	113	2.00E-24	225	76	33
AF529084	780	121	4.00E-27	260	83	31	U76543	1023	107	6.00E-23	187	70	37
L81176	5735	119	2.00E-26	195	77	39	AF003905	1023	113	2.00E-24	225	78	33
AY278546	607	119	2.00E-26	195	77	39	AF003905	1023	105	3.00E-22	187	69	36
AY278537	599	119	2.00E-26	195	77	39	AB018711	1194	113	2.00E-24	215	86	40
X98462	1301	119	2.00E-26	197	78	39	AB018711	1194	55.8	3.00E-07	130	43	33
X98462	1301	92.4	3.00E-18	309	90	29	X98464	1019	112	2.00E-24	225	76	33
AF416443	388	118	4.00E-26	117	60	51	X98464	1019	105	4.00E-22	188	89	37
X98280	1301	118	4.00E-26	196	78	39	AF016232	1302	112	2.00E-24	196	73	37
X98280	1301	98.6	4.00E-20	326	91	27	AF016232	1302	102	3.00E-21	327	93	28
AF416435	381	118	5.00E-26	120	59	49	X98461	1301	112	3.00E-24	198	75	38
AB181529	1701	118	5.00E-26	180	68	37	X98461	1301	95.5	3.00E-19	326	90	27
AB181529	1701	75.9	3.00E-13	84	34	40	AB018724	510	111	4.00E-24	125	61	48
AB018719	1449	118	5.00E-26	254	87	34	AB018722	450	111	4.00E-24	125	61	48
AB018719	1449	58.5	4.00E-08	221	60	27	X98281	1301	110	7.00E-24	195	74	37
AF416449	395	117	8.00E-26	117	59	50	X98281	1301	89	3.00E-17	321	85	26
AF416448	399	117	8.00E-26	117	59	50	AB018714	528	110	7.00E-24	174	74	42
AF416447	399	117	8.00E-26	117	59	50	AB181534	1704	110	1.00E-23	180	64	35
AF416446	394	117	8.00E-26	117	59	50	AB181534	1704	75.5	3.00E-13	83	34	40
AF416445	396	117	8.00E-26	117	59	50	BX571662	349859	109	2.00E-23	294	89	30
AF416444	399	117	8.00E-26	117	59	50	BX571662	349859	73.2	2.00E-12	122	44	36
AF416442	389	117	8.00E-26	117	59	50	AB181533	1704	109	2.00E-23	180	64	35
AF416441	397	117	8.00E-26	117	59	50	AB181533	1704	75.5	3.00E-13	83	34	40
AF416440	393	117	8.00E-26	117	59	50	AB181526	1704	109	2.00E-23	180	63	35
AF416439	393	117	8.00E-26	117	59	50	AB181526	1704	74.7	6.00E-13	83	34	40
AF416438	396	117	8.00E-26	117	59	50	AF290503	1728	109	2.00E-23	184	63	34
AF416436	399	117	8.00E-26	117	59	50	AF290503	1728	79	3.00E-14	107	43	40
AF416434	399	117	8.00E-26	117	59	50	AF290502	1728	109	2.00E-23	184	63	34
AF416433	397	117	8.00E-26	117	59	50	AF290502	1728	79	3.00E-14	107	43	40
AB181536	1698	117	1.00E-25	180	68	37	AF290501	1728	109	2.00E-23	184	63	34
AB181536	1698	75.5	3.00E-13	83	34	40	AF290501	1728	79	3.00E-14	107	43	40
AB181530	1701	117	1.00E-25	180	68	37	AF290500	1722	109	2.00E-23	184	63	34
AB181530	1701	74.7	6.00E-13	83	34	40	AF290500	1722	79.3	2.00E-14	124	47	37
AB181528	1701	117	1.00E-25	180	68	37	AF290499	1722	109	2.00E-23	184	63	34
AB181528	1701	74.7	6.00E-13	83	34	40	AF290498	1722	79.3	2.00E-14	124	47	37
AB181524	1701	117	1.00E-25	180	68	37	AF290498	1722	109	2.00E-23	184	63	34
AB181524	1701	74.7	6.00E-13	83	34	40	AF290498	1722	79.3	2.00E-14	124	47	37
AB181522	1698	117	1.00E-25	180	68	37	AF290497	1722	109	2.00E-23	184	63	34
AB181522	1698	75.5	3.00E-13	83	34	40	AF290497	1722	79.3	2.00E-14	124	47	37
AB181521	1698	117	1.00E-25	180	68	37	AF290496	1722	109	2.00E-23	184	63	34
AB181521	1698	75.5	3.00E-13	83	34	40	AF290496	1722	79.3	2.00E-14	124	47	37
AB105426	3847	115	1.00E-25	132	58	43	AF202168	7756	109	2.00E-23	184	63	34
AB105426	3847	102	2.00E-21	297	90	30	AF202168	7756	105	3.00E-22	184	63	34
AJ277354	663	115	1.00E-25	238	74	31	AF202168	7756	77	1.00E-13	83	37	44
AB181523	1701	115	1.00E-25	180	68	37	AF202168	7756	77	1.00E-13	83	37	44
AB181523	1701	74.7	6.00E-13	83	34	40	AF202168	7756	73.6	1.00E-12	83	34	40
AB181525	1689	116	2.00E-25	180	67	37	AF202168	7756	63.5	1.00E-09	133	36	27
AB181525	1689	76.6	2.00E-13	173	50	28	AF202168	7756	47	1.00E-04	52	26	50
AB181532	1695	115	2.00E-25	180	67	37	AF050191	1731	109	2.00E-23	184	62	33
AB181532	1695	78.2	5.00E-14	127	45	35	AF050191	1731	77.4	9.00E-14	156	52	33
AF016231	1302	115	1.00E-25	196	76	38	AJ297532	1370	108	3.00E-23	144	65	45
AF016231	1302	103	1.00E-21	327	93	28	AJ297532	1370	98.6	4.00E-20	275	84	30
AF016230	1302	115	2.00E-25	196	76	38	AB181531	1704	108	4.00E-23	180	63	35
AF016230	1302	102	3.00E-21	327	93	28	AB181531	1704	75.5	3.00E-13	83	34	40
AF016229	1302	115	2.00E-25	196	76	38	AB098070	1713	108	4.00E-23	181	66	36
AF016229	1302	102	3.00E-21	327	93	28	AB098070	1713	72	4.00E-12	83	34	40
AF003906	1004	115	2.00E-25	265	87	32	AB098067	1725	108	4.00E-23	184	63	34
AF003906	1004	110	1.00E-23	174	66	37	AB098067	1725	77.4	9.00E-14	159	55	34
AB181527	1698	115	3.00E-25	180	67	37	AF140252	2031	108	4.00E-23	184	63	34
AB181527	1698	75.5	3.00E-13	83	34	40	AF140252	2031	77	1.00E-13	83	37	44
AF416437	379	114	5.00E-25	127	61	48	AF270499	1146	108	4.00E-23	215	72	33
AY024344	783	114	7.00E-25	257	80	31	AF050195	1731	108	4.00E-23	184	62	33
X98465	2099	114	7.00E-25	301	93	30	AF050195	1731	77.4	9.00E-14	83	37	44
X98465	2099	99.8	2.00E-20	297	91	30	AF050190	1731	108	4.00E-23	184	63	34
AJ297533	1319	113	1.00E-24	237	83	35	AF050190	1731	77.4	9.00E-14	124	44	35
AJ297533	1319	103	1.00E-21	254	79	31	AF050184	1731	108	4.00E-23	184	62	33
AB181535	1713	113	2.00E-24	180	66	36	AF050184	1731	77	1.00E-13	83	37	44
AB181535	1713	74.7	6.00E-13	83	33	39	AB070578	1458	108	4.00E-23	183	67	36

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AB070578	1458	80.1	1.00E-14	83	38	45	U28497	1055	105	3.00E-22	308	87	28
M64671	4200	108	4.00E-23	184	63	34	AB018726	372	105	3.00E-22	124	57	45
M64671	4200	105	3.00E-22	184	64	34	M35141	1932	105	3.00E-22	184	64	34
M64671	4200	79.7	2.00E-14	124	46	37	M35141	1932	72.8	2.00E-12	124	43	34
M64671	4200	73.9	1.00E-12	124	43	34	AY102622	7647	105	4.00E-22	184	62	33
J05635	3832	108	4.00E-23	184	62	33	AY102622	7647	73.6	1.00E-12	83	34	40
J05635	3832	105	3.00E-22	184	63	34	AY278540	550	105	4.00E-22	186	69	37
J05635	3832	77	1.00E-13	83	37	44	AE017149	266956	104	5.00E-22	188	67	35
J05635	3832	71.6	5.00E-12	83	33	39	AE017149	266956	97.8	7.00E-20	170	60	35
M64670	4200	108	4.00E-23	184	63	34	AE017149	266956	76.3	2.00E-13	297	74	24
M64670	4200	105	3.00E-22	184	64	34	AEC17149	266956	68.9	3.00E-11	115	40	34
M64670	4200	79.7	2.00E-14	124	46	37	AB103059	1461	104	5.00E-22	183	65	35
M64670	4200	73.9	1.00E-12	124	43	34	AB103059	1461	80.1	1.00E-14	83	38	45
AE000533	11152	108	5.00E-23	187	67	35	AB103056	1461	104	5.00E-22	183	65	35
AE000533	11152	77.4	9.00E-14	121	47	38	AB103056	1461	79	3.00E-14	83	37	44
AB073915	1458	108	5.00E-23	183	66	36	AB103053	1461	104	5.00E-22	183	65	35
AB073915	1458	80.1	1.00E-14	83	38	45	AB103053	1461	79	3.00E-14	83	37	44
AY304577	1545	107	6.00E-23	166	62	37	AB073918	1473	104	5.00E-22	180	63	35
AY304577	1545	77.4	9.00E-14	121	47	38	AB073918	1473	81.6	5.00E-15	174	54	31
AB098069	1722	107	6.00E-23	181	65	35	L08908	1800	104	5.00E-22	189	63	33
AB098069	1722	82	4.00E-15	162	56	34	L08908	1800	80.5	1.00E-14	123	48	39
AB098068	1722	107	6.00E-23	181	65	35	AE005820	14021	104	7.00E-22	297	76	25
AB098068	1722	79	3.00E-14	83	37	44	AE005820	14021	99.4	2.00E-20	298	76	25
AF050186	1728	107	6.00E-23	181	65	35	AE005820	14021	99	3.00E-20	292	75	25
AF050186	1728	82	4.00E-15	162	56	34	AB084912	1461	104	7.00E-22	183	64	34
AY714226	1545	107	8.00E-23	187	67	35	AB084912	1461	82.8	2.00E-15	170	56	32
AY714226	1545	77.4	9.00E-14	121	47	38	AB084911	1461	104	7.00E-22	183	64	34
AY319299	1545	107	8.00E-23	187	67	35	AB084911	1461	82.8	2.00E-15	170	56	32
AY319299	1545	77.4	9.00E-14	121	47	38	AF060185	1731	104	7.00E-22	181	64	35
AF479024	1545	107	8.00E-23	187	67	35	AF060185	1731	79.3	2.00E-14	121	44	36
AF479024	1545	77.4	9.00E-14	121	47	38	AB080202	1473	104	7.00E-22	181	63	34
Z29327	3879	107	8.00E-23	182	67	36	AB080202	1473	83.2	2.00E-15	174	55	31
Z29327	3879	103	9.00E-22	182	69	37	AB103052	1473	103	9.00E-22	184	63	34
Z29327	3879	79.3	2.00E-14	107	43	40	AB103052	1473	82.8	2.00E-15	174	59	33
Z29327	3879	73.9	1.00E-12	107	40	37	AF050194	1728	103	9.00E-22	171	62	36
Y11602	1900	107	8.00E-23	249	79	31	AF050194	1728	78.6	4.00E-14	107	43	40
Y11602	1900	75.1	5.00E-13	148	50	33	BX571661	346613	103	1.00E-21	192	63	32
AY155232	1545	107	8.00E-23	187	67	35	BX571661	346613	94.7	6.00E-19	333	85	25
AY155232	1545	77.4	9.00E-14	121	47	38	Y11762	4057	103	1.00E-21	181	64	35
AB103061	1725	107	8.00E-23	171	64	37	Y11762	4057	102	3.00E-21	183	65	35
AB103061	1725	79.7	2.00E-14	121	44	36	Y11762	4057	79.3	2.00E-14	148	50	33
M74578	1800	107	8.00E-23	182	67	36	Y11762	4057	71.2	7.00E-12	105	38	36
M74578	1800	73.9	1.00E-12	107	40	37	AF060193	1719	103	1.00E-21	181	67	37
AE001449	13631	107	8.00E-23	187	67	35	AF060193	1719	72.8	2.00E-12	132	45	34
AE001449	13631	77.4	9.00E-14	121	47	38	M82917	2236	103	1.00E-21	192	63	32
L08907	1800	107	8.00E-23	187	67	35	M82917	2236	94.7	6.00E-19	333	85	25
L08907	1800	77.4	9.00E-14	121	47	38	AB073917	1461	103	1.00E-21	183	63	34
X57173	1731	107	1.00E-22	181	65	35	AB073917	1461	80.1	1.00E-14	83	38	45
X57173	1731	82	4.00E-15	162	56	34	AB103055	1473	103	2.00E-21	219	68	31
AF050197	1719	107	1.00E-22	171	64	37	AB103055	1473	84	1.00E-15	171	56	32
AF050197	1719	77.8	7.00E-14	132	48	36	AF08835	2310	103	2.00E-21	297	75	25
AF050196	1719	107	1.00E-22	181	65	35	AF08835	2310	99	3.00E-20	292	75	25
AF050196	1719	78.2	5.00E-14	132	48	36	AE017138	290924	102	2.00E-21	223	67	30
AF050192	1719	107	1.00E-22	181	65	35	AE017138	290924	47.8	8.00E-05	66	22	33
AF050192	1719	79.3	2.00E-14	132	48	36	AE013947	10446	102	2.00E-21	223	67	30
AF050188	1719	107	1.00E-22	181	65	35	AE013947	10446	57.4	1.00E-07	84	28	33
AF050188	1719	78.6	4.00E-14	83	37	44	AJ414144	208050	102	2.00E-21	223	67	30
AB103060	1716	106	1.00E-22	184	61	33	AJ414144	208050	95.9	2.00E-19	295	76	25
AB103080	1716	79.7	2.00E-14	124	46	37	AJ414144	208050	92	4.00E-18	309	79	25
AL139078	263335	106	2.00E-22	171	63	36	AY751741	1630	102	3.00E-21	169	61	36
AL139078	263335	102	3.00E-21	181	66	36	AY751741	1630	62	4.00E-09	141	42	29
AL139078	263335	79.3	2.00E-14	132	48	36	AB103054	1470	102	3.00E-21	219	68	31
AL139078	263335	72.8	2.00E-12	132	45	34	AB103054	1470	83.6	1.00E-15	171	59	34
AB018730	375	106	2.00E-22	125	57	45	AB018720	360	102	3.00E-21	119	55	46
AB018728	393	106	2.00E-22	125	57	45	AF369587	1671	101	5.00E-21	178	59	33
AB103058	1461	105	2.00E-22	183	64	34	AF369587	1671	70.1	1.00E-11	113	41	36
AB103058	1461	80.1	1.00E-14	83	38	45	AF050187	1728	101	5.00E-21	181	65	35
AB018716	495	105	2.00E-22	165	68	41	AF050187	1728	77.8	7.00E-14	296	78	26

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
Y11601	1900	101	6.00E-21	240	73	30	AY319298	1533	82	4.00E-15	311	78	25
Y11601	1900	77.8	7.00E-14	329	89	27	AY278532	503	94.7	6.00E-19	164	59	35
AB103051	1476	101	6.00E-21	174	61	35	Z25773	404	93.6	1.00E-18	154	59	38
AB103051	1476	82.8	2.00E-15	175	58	33	BX294145	300350	92.8	2.00E-18	169	62	36
AF369581	1671	101	6.00E-21	178	59	33	BX294145	300350	85.5	3.00E-16	126	55	43
AF369581	1671	70.1	1.00E-11	113	41	36	AJ297534	1512	92.4	3.00E-18	208	66	31
AF050189	1719	101	6.00E-21	181	66	36	AJ297534	1512	66.2	2.00E-10	252	61	24
AF050189	1719	71.6	5.00E-12	132	45	34	X96435	3919	91.3	6.00E-18	313	77	24
AE000767	13413	100	8.00E-21	209	62	29	X96435	3919	79.7	2.00E-14	313	75	23
AE000767	13413	62	4.00E-09	129	39	30	X96435	3919	71.6	5.00E-12	324	75	23
AB103050	1500	100	8.00E-21	153	61	39	AE009023	9808	91.3	6.00E-18	313	77	24
AB103050	1500	85.9	3.00E-16	316	82	25	AE009023	9808	79.7	2.00E-14	313	75	23
AF369584	1668	100	1.00E-20	175	62	35	AE009023	9808	71.6	5.00E-12	324	75	23
AF369584	1668	71.6	5.00E-12	287	73	25	AE007989	10029	91.3	6.00E-18	313	77	24
AF369583	1574	100	1.00E-20	175	62	35	AE007989	10029	79.7	2.00E-14	313	75	23
AF369583	1574	74.7	6.00E-13	287	72	25	AE007989	10029	71.6	5.00E-12	324	75	23
AF369580	582	100	1.00E-20	175	62	35	U95165	21846	91.3	6.00E-18	313	77	24
AF369586	1682	100	1.00E-20	175	61	34	U95165	21846	79.7	2.00E-14	313	75	23
AF369586	1682	68.6	4.00E-11	74	31	41	U95165	21846	71.6	5.00E-12	324	75	23
AF369585	1662	100	1.00E-20	175	61	34	U95165	21846	53.5	1.00E-06	256	60	23
AF369585	1662	68.6	4.00E-11	74	31	41	U95165	21846	46.2	2.00E-04	140	34	24
AF369582	1682	100	1.00E-20	175	61	34	X80701	4423	91.3	6.00E-18	313	77	24
AF369582	1682	68.6	4.00E-11	74	31	41	X80701	4423	79.7	2.00E-14	313	75	23
AF369579	582	100	1.00E-20	175	61	34	X80701	4423	71.6	5.00E-12	324	75	23
AF369578	582	100	1.00E-20	175	61	34	AB110835	909	90.5	1.00E-17	301	81	28
AF369577	582	100	1.00E-20	175	61	34	L38478	1446	90.5	1.00E-17	150	51	34
AB018712	345	99.8	2.00E-20	115	58	50	U17575	2803	90.5	1.00E-17	200	57	28
AB073916	1476	99.8	2.00E-20	174	59	33	U17575	2803	56.6	2.00E-07	146	47	32
AB073916	1476	85.9	3.00E-16	172	57	33	AE001699	13774	89.7	2.00E-17	248	63	25
AE005755	12263	98	3.00E-20	292	72	24	AE0142921207381	88.6	4.00E-17	298	76	25	
AE005755	12263	98.6	4.00E-20	292	72	24	U42432	305	88.6	4.00E-17	87	45	51
AE005755	12263	97.4	9.00E-20	292	70	23	AF019251	3817	88.6	4.00E-17	298	76	25
M26945	1719	99	3.00E-20	171	58	33	AY751740	1595	87.4	9.00E-17	142	52	36
M26945	1719	79.7	2.00E-14	124	46	37	AY751740	1595	63.9	1.00E-09	94	34	36
AF040258	3447	99	3.00E-20	292	72	24	AE009654	10807	86.7	2.00E-16	298	75	25
AF040258	3447	98.6	4.00E-20	292	72	24	BA0000127036071	84.7	6.00E-16	329	75	22	
AF040268	3447	95.5	3.00E-19	292	70	23	BA0000127036071	59.7	2.00E-08	356	80	22	
AE013020	10181	98.2	5.00E-20	290	83	28	AB110836	909	84.3	7.00E-16	92	44	47
BX294139	287650	98.2	5.00E-20	284	88	30	U42431	305	84	1.00E-15	87	43	49
BX294139	287650	81.3	6.00E-15	143	57	39	JO1556	1193	82.4	3.00E-15	298	70	23
AB018718	1191	98.2	5.00E-20	138	67	35	AJ418317	1384	79	3.00E-14	149	49	32
AB018718	1191	51.2	7.00E-06	91	27	29	AJ418317	1384	63.9	1.00E-09	109	39	35
AE017148	317022	97.8	7.00E-20	170	60	35	AJ418316	1384	79	3.00E-14	149	49	32
AE017148	317022	76.3	2.00E-13	297	74	24	AJ418316	1384	63.9	1.00E-09	109	39	35
BX294140	307050	97.4	9.00E-20	257	76	29	AJ418322	1376	77.8	7.00E-14	136	45	33
BX294140	307050	79.3	2.00E-14	250	70	28	AJ418322	1376	67.8	7.00E-11	294	70	23
BX294140	307050	34.3	0.89	120	34	28	AJ418321	1376	77.8	7.00E-14	136	45	33
BX294140	307050	33.1	2	116	34	29	AJ418321	1376	65.9	3.00E-10	294	69	23
X60746	1800	97.1	1.00E-19	170	57	33	AJ418320	1377	77.8	7.00E-14	136	45	33
X60746	1800	82.4	3.00E-15	311	78	25	AJ418320	1377	65.9	3.00E-10	294	69	23
AB103057	1500	97.1	1.00E-19	222	73	32	AJ418319	1377	77.8	7.00E-14	136	45	33
AB103057	1500	82.4	3.00E-15	174	55	31	AJ418319	1377	65.9	3.00E-10	294	69	23
AY304576	1533	96.3	2.00E-19	170	57	33	AJ418318	1377	77.8	7.00E-14	136	45	33
AY304576	1533	83.2	2.00E-15	311	79	25	AJ418318	1377	65.9	3.00E-10	294	69	23
AY714225	1533	96.3	2.00E-19	170	57	33	AY134860	420	77.4	9.00E-14	74	38	51
AY714225	1533	82	4.00E-15	311	78	25	AJ297531	1299	77.4	9.00E-14	315	74	23
AJ297530	1280	96.3	2.00E-19	251	76	30	AB035615	600	77.4	9.00E-14	134	46	34
AJ297530	1280	91.7	5.00E-18	270	73	27	AB035614	600	77.4	9.00E-14	134	46	34
AY155231	1533	96.3	2.00E-19	170	57	33	AB035613	600	77.4	9.00E-14	134	46	34
AY155231	1533	82.4	3.00E-15	311	78	25	AF398973	437	77	1.00E-13	158	53	33
AE000574	10042	96.3	2.00E-19	170	57	33	AB091714	600	77	1.00E-13	134	46	34
AE000574	10042	82.4	3.00E-15	311	78	25	AB091713	600	77	1.00E-13	134	46	34
AE001487	17223	96.3	2.00E-19	170	57	33	AB091712	600	77	1.00E-13	134	46	34
AE001487	17223	82.4	3.00E-15	311	78	25	AB091711	600	77	1.00E-13	134	46	34
L36137	1548	96.3	2.00E-19	172	69	40	AB091710	600	77	1.00E-13	134	46	34
L36137	1548	45.8	3.00E-04	69	33	47	AB091709	600	77	1.00E-13	134	46	34
L36137	1548	45.4	4.00E-04	60	31	51	AB091708	600	77	1.00E-13	134	46	34
AY319298	1533	96.9	2.00E-19	170	57	33	AB091707	600	77	1.00E-13	134	46	34

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AB091706	600	77	1.00E-13	134	46	34	AB178325	597	73.9	1.00E-12	133	44	33
AB091706	600	77	1.00E-13	134	46	34	AY278543	479	73.2	2.00E-12	162	54	33
AB091704	600	77	1.00E-13	134	46	34	AB189459	597	72	4.00E-12	133	43	32
AB091703	600	77	1.00E-13	134	46	34	CR354386	888	71.6	5.00E-12	294	67	22
AB091702	600	77	1.00E-13	134	46	34	AF072133	582	71.2	7.00E-12	100	41	41
AB091701	600	77	1.00E-13	134	46	34	AY278539	474	71.2	7.00E-12	160	53	33
AB091700	600	77	1.00E-13	134	46	34	AY278536	463	71.2	7.00E-12	160	53	33
AB037130	600	77	1.00E-13	134	46	34	AY278544	468	69.7	2.00E-11	159	52	32
AB037129	600	77	1.00E-13	134	46	34	CR354387	852	69.7	2.00E-11	284	61	21
AB037128	600	77	1.00E-13	134	46	34	AF030241	751	68.9	3.00E-11	186	50	26
AB037127	600	77	1.00E-13	134	46	34	AF030241	751	38.5	0.047	49	20	40
AY505350	209	76.5	2.00E-13	69	39	56	U85322	1105	68.9	3.00E-11	288	68	23
AB052665	600	75.9	3.00E-13	134	45	33	AL139076	317511	68.6	4.00E-11	288	66	22
AB035621	600	75.9	3.00E-13	134	45	33	AF461538	7699	68.2	6.00E-11	257	63	24
AB035620	600	75.9	3.00E-13	134	45	33	AY278541	466	67.4	9.00E-11	158	52	32
AB035619	600	75.9	3.00E-13	134	45	33	U26705	641	67.4	9.00E-11	221	58	26
AB035618	600	75.9	3.00E-13	134	45	33	U26704	641	67.4	9.00E-11	221	58	26
AB035617	600	75.9	3.00E-13	134	45	33	AF398975	351	57	1.00E-10	129	46	35
AB035616	600	75.9	3.00E-13	134	45	33	AF398974	351	67	1.00E-10	129	46	35
AB035612	600	75.9	3.00E-13	134	45	33	AE017253	302836	66.6	2.00E-10	285	65	22
AB035611	600	75.9	3.00E-13	134	45	33	V01370	1149	66.2	2.00E-10	37	34	91
AB035610	600	75.9	3.00E-13	134	45	33	AF398982	378	65.1	5.00E-10	130	41	31
AB035609	600	75.9	3.00E-13	134	45	33	AF398981	378	65.1	5.00E-10	130	41	31
AB035608	600	75.9	3.00E-13	134	45	33	AF398976	378	65.1	5.00E-10	130	42	32
AB035607	600	75.9	3.00E-13	134	45	33	AE001695	13302	65.1	5.00E-10	258	62	24
AB035606	600	75.9	3.00E-13	134	45	33	AF030240	766	63.5	1.00E-09	202	51	25
AB035605	600	75.9	3.00E-13	134	45	33	AF030240	766	38.5	0.047	49	20	40
AB035604	600	75.9	3.00E-13	134	45	33	AE001240	14244	63.2	2.00E-09	268	60	22
AB035603	600	75.9	3.00E-13	134	45	33	AY342020	584	63.2	2.00E-09	190	50	26
AB035602	600	75.9	3.00E-13	134	45	33	AL591784	300000	62.4	3.00E-09	321	68	21
AB035601	600	75.9	3.00E-13	134	45	33	AL591784	300000	50.4	1.00E-05	87	29	35
AB035600	600	75.9	3.00E-13	134	45	33	AL591784	300000	45.8	3.00E-04	245	56	22
AB035599	600	75.9	3.00E-13	134	45	33	AL591784	300000	40	0.016	87	24	27
AB035598	600	75.9	3.00E-13	134	45	33	AJ297535	949	62.4	3.00E-09	141	46	32
AB035597	600	75.9	3.00E-13	134	45	33	AJ297535	949	40.4	1.00E-04	57	26	45
AB035596	600	75.9	3.00E-13	134	45	33	AJ297536	949	31.2	1.00E-04	78	21	26
AB035595	600	75.9	3.00E-13	134	45	33	L49337	19824	62	4.00E-09	321	65	20
AB091808	597	75.5	3.00E-13	133	45	33	L49337	19824	50.4	1.00E-05	149	43	28
AB091806	597	75.5	3.00E-13	133	45	33	L49337	19824	45.1	5.00E-04	146	38	26
AB178780	597	75.5	3.00E-13	133	45	33	L49337	19824	40	0.016	87	24	27
AB178779	597	75.5	3.00E-13	133	45	33	AF264897	608	61.2	7.00E-09	208	52	25
AB178335	597	75.5	3.00E-13	133	45	33	AF264883	608	61.2	7.00E-09	208	52	25
AB178334	597	75.5	3.00E-13	133	45	33	AY342021	584	61.2	7.00E-09	190	49	26
AB091812	597	75.1	5.00E-13	133	45	33	AY342019	584	61.2	7.00E-09	190	48	25
AB091811	597	75.1	5.00E-13	133	45	33	AF264899	608	60.8	9.00E-09	208	51	24
AB091810	597	75.1	5.00E-13	133	45	33	AF264898	608	60.8	9.00E-09	208	51	24
AB091809	597	75.1	5.00E-13	133	45	33	AF264896	608	60.5	1.00E-08	208	50	24
AB091805	597	75.1	5.00E-13	133	45	33	AF264895	608	60.5	1.00E-08	208	50	24
AB178333	597	75.1	5.00E-13	133	45	33	AF264894	608	60.5	1.00E-08	208	50	24
AB091715	600	74.7	6.00E-13	134	45	33	AF264893	608	60.5	1.00E-08	208	50	24
BX571669	349970	74.3	8.00E-13	278	65	23	AF264892	608	60.5	1.00E-08	208	50	24
AB109246	596	74.3	8.00E-13	133	44	33	AF264891	608	60.5	1.00E-08	208	50	24
AB109245	596	74.3	8.00E-13	133	44	33	AF264890	608	60.5	1.00E-08	208	50	24
AB109244	596	74.3	8.00E-13	133	44	33	AF264888	608	60.5	1.00E-08	208	50	24
AB109243	596	74.3	8.00E-13	133	44	33	AF264887	608	60.5	1.00E-08	208	50	24
AB109242	596	74.3	8.00E-13	133	44	33	AY342027	584	60.5	1.00E-08	190	48	25
AB109241	596	74.3	8.00E-13	133	44	33	AY342026	584	60.5	1.00E-08	190	48	25
AB091814	597	73.9	1.00E-12	133	44	33	AY342026	584	60.5	1.00E-08	190	48	25
AB091813	597	73.9	1.00E-12	133	44	33	AY342024	584	60.5	1.00E-08	190	48	25
AB091807	597	73.9	1.00E-12	133	44	33	AY342023	584	60.5	1.00E-08	190	48	25
AB180460	597	73.9	1.00E-12	133	44	33	AY342022	584	60.5	1.00E-08	190	48	25
AB178331	597	73.9	1.00E-12	133	44	33	AY342018	584	60.5	1.00E-08	190	48	25
AB178332	597	73.9	1.00E-12	133	44	33	AB001703	584	60.5	1.00E-08	190	48	25
AB178330	597	73.9	1.00E-12	133	44	33	AB001701	584	60.5	1.00E-08	190	48	25
AB178329	597	73.9	1.00E-12	133	44	33	AB001700	584	60.5	1.00E-08	190	48	25
AB178328	597	73.9	1.00E-12	133	44	33	AF398984	378	60.1	2.00E-08	124	41	33
AB178327	597	73.9	1.00E-12	133	44	33	AF398983	378	60.1	2.00E-08	124	41	33
AB178326	597	73.9	1.00E-12	133	44	33	AF264886	608	60.1	2.00E-08	208	49	23

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AE007721	8198	60.1	2.00E-08	289	65	22	AY366639	426	47	1.00E-04	132	36	27
AE007721	8198	42.4	0.003	310	69	22	AE004539	25022	46.6	2.00E-04	134	34	25
AF264889	608	60.1	2.00E-08	208	49	23	AY007305	1233	45.6	2.00E-04	116	29	25
AF264885	608	60.1	2.00E-08	208	49	23	AF408410	423	46.2	2.00E-04	126	35	27
AF264882	608	60.1	2.00E-08	208	49	23	AJ311995	235	46.2	2.00E-04	71	25	35
AF264881	608	60.1	2.00E-08	208	49	23	AJ311997	235	46.2	2.00E-04	71	25	35
AF264880	608	60.1	2.00E-08	208	49	23	AJ311995	235	46.2	2.00E-04	71	25	35
AF264879	608	60.1	2.00E-08	208	49	23	AJ311995	235	46.2	2.00E-04	71	25	35
AF396980	378	59.7	2.00E-08	138	44	31	AJ311993	235	46.2	2.00E-04	71	25	35
AF396979	378	59.7	2.00E-08	138	44	31	AE001532	11113	46.2	2.00E-04	149	40	26
AF396978	378	59.7	2.00E-08	138	44	31	AF015108	267	46.2	2.00E-04	77	21	27
AF396977	378	59.7	2.00E-08	138	44	31	Y15098	477	45.8	3.00E-04	139	33	23
AB001702	584	59.7	2.00E-08	190	48	25	Y15095	463	45.8	3.00E-04	139	33	23
AF264884	608	59.3	3.00E-08	208	49	23	Y15094	481	45.8	3.00E-04	139	33	23
AF264901	602	58.5	4.00E-08	206	50	24	Y15091	463	45.8	3.00E-04	139	33	23
AB018725	537	58.5	4.00E-08	69	29	42	AF015109	267	45.8	3.00E-04	77	21	27
AB018723	408	58.5	4.00E-08	69	29	42	AY365212	402	45.4	4.00E-04	132	35	26
M21445	867	58.5	4.00E-08	46	28	60	Y15101	490	45.4	4.00E-04	139	33	23
M21445	867	41.6	0.006	307	79	25	Y15099	466	45.4	4.00E-04	139	33	23
AY083505	621	58.2	6.00E-08	161	41	25	Y15097	479	45.4	4.00E-04	139	33	23
AY083504	621	58.2	6.00E-08	161	41	25	Y15092	463	45.4	4.00E-04	139	33	23
AJ297537	384	57.8	8.00E-08	64	28	43	Y15090	473	45.4	4.00E-04	139	33	23
AF264900	608	57.8	8.00E-08	208	51	24	Y15089	478	45.4	4.00E-04	139	33	23
AY450560	570	57.4	1.00E-07	185	47	25	AF116904	467	45.4	4.00E-04	143	36	25
AY278538	432	57	1.00E-07	149	47	31	Y15100	456	45.1	5.00E-04	139	33	23
AY278533	417	57	1.00E-07	135	41	30	X89239	1310	45.1	5.00E-04	105	29	27
U62056	2236	56.2	2.00E-07	47	27	57	AF228032	476	45.1	5.00E-04	146	37	25
AJ297536	340	52.8	2.00E-07	43	27	62	D12510	832	45.1	5.00E-04	24	23	95
AJ297536	340	28.5	2.00E-07	31	13	41	AY228590	321	44.7	7.00E-04	94	24	25
AF322547	19758	55.8	3.00E-07	132	36	27	BX5719663173005	44.7	7.00E-04	292	64	21	
M34710	684	55.3	3.00E-07	63	27	42	BX5719663173005	42.7	0.002	320	69	21	
AJ251711	1260	55.1	5.00E-07	193	48	24	BX5719663173005	40.4	0.012	278	59	21	
AJ251711	1260	39.7	0.021	72	22	30	BX5719663173005	40.4	0.012	291	65	22	
AB018721	282	55.1	5.00E-07	49	23	46	BX5719663173005	39.3	0.028	302	59	19	
AE011584	11491	54.7	6.00E-07	110	34	30	AF228034	476	44.7	7.00E-04	146	37	25
AY083503	615	54.7	6.00E-07	156	41	25	AF228033	476	44.7	7.00E-04	146	37	25
AY083502	615	54.7	6.00E-07	156	41	25	AF354560	321	44.7	7.00E-04	94	24	25
AE017300	358408	54.7	6.00E-07	110	34	30	AF354559	321	44.7	7.00E-04	94	24	25
BX640415	347071	54.7	6.00E-07	136	43	31	AF354558	321	44.7	7.00E-04	94	24	25
Y15093	506	53.9	1.00E-06	159	39	24	AF355599	321	44.7	7.00E-04	94	24	25
AB018731	363	53.9	1.00E-06	92	32	34	Y15095	480	44.3	9.00E-04	139	33	23
AB018729	363	53.9	1.00E-06	92	32	34	AE006319	10302	44.3	9.00E-04	287	50	17
AB018727	369	53.9	1.00E-06	92	32	34	AF119150	18505	44.3	9.00E-04	228	52	22
AE009025	11897	53.5	1.00E-06	256	60	23	M33808	852	44.3	9.00E-04	30	21	70
AE009025	11897	46.2	2.00E-04	140	34	24	M33808	852	37.4	0.1	31	18	58
AE007991	9626	53.5	1.00E-06	256	60	23	AE004223	24353	43.9	0.001	228	52	22
AE007991	9626	45.2	2.00E-04	140	34	24	AF354561	321	43.9	0.001	94	24	25
AB018713	369	53.5	1.00E-06	108	33	30	AF015104	267	43.9	0.001	77	21	27
AF515473	528	53.1	2.00E-06	92	37	40	AY226588	321	43.5	0.001	91	28	30
AF515473	528	49.7	2.00E-05	132	41	31	AY226587	321	43.5	0.001	91	28	30
AY357714	479	52.4	3.00E-06	150	40	26	AF354554	321	43.5	0.001	91	28	30
AY362359	436	52	4.00E-06	144	38	26	AF354553	321	43.5	0.001	91	28	30
Y15088	506	52	4.00E-06	159	38	23	AF354548	321	43.5	0.001	91	28	30
U26679	1573	52	4.00E-06	108	32	29	AF015105	267	43.5	0.001	77	20	25
AF497995	544	51.6	5.00E-06	181	44	24	AF015101	267	43.5	0.001	77	20	25
AB018715	468	51.6	5.00E-06	88	27	30	AF015097	267	43.5	0.001	77	20	25
BX572605	349746	50.8	9.00E-06	284	69	24	AF015089	267	43.5	0.001	77	20	25
BX572605	349746	44.7	7.00E-04	194	49	25	AY226589	321	43.1	0.002	91	27	29
BX572600	349640	50.8	9.00E-06	284	70	24	X51740	1766	43.1	0.002	21	21	100
M57565	3608	50.4	1.00E-05	87	29	33	BA0000381857073	43.1	0.002	289	63	21	
M57565	3608	45.8	3.00E-04	245	56	22	BA0000381857073	33.1	2	120	30	25	
M24526	3530	50.4	1.00E-05	149	43	28	AF354555	321	43.1	0.002	91	27	29
M24526	3530	42	0.004	87	23	26	AF354550	321	43.1	0.002	91	27	29
AY533375	7214	49.3	3.00E-05	134	34	25	AF354549	321	43.1	0.002	91	27	29
AY533375	7214	48.5	5.00E-05	130	34	26	AF354547	321	43.1	0.002	91	27	29
AB018717	234	48.9	3.00E-05	74	28	37	AF354546	321	43.1	0.002	91	27	29
D26168	866	47.8	8.00E-05	24	24	100	X89238	1310	42.7	0.002	99	27	27
D26167	832	47.8	8.00E-05	24	24	100	AF354552	321	42.7	0.002	81	26	32

Fig. 25A-Continued

Hit_ID	Length	BitScore	Expected	Length	#Ident	%S	Hit_ID	Length	BitScore	Expected	Length	#Ident	%S
AF354551	321	42.7	0.002	81	26	32	BA0000332820462	34.3	0.89	299	60	20	
AF015106	267	42.7	0.002	77	20	25	U66234	390	39.3	0.028	97	26	26
BX49358	348408	42.4	0.003	169	38	22	AL596163	231450	38.9	0.036	261	57	21
BX571658	346792	42.4	0.003	168	37	22	U25727	108	38.9	0.036	34	21	61
AL591982	295050	42.4	0.003	273	58	21	AY708387	1232	38.5	0.047	260	55	21
AF354557	321	42.4	0.003	81	25	30	AY708385	1250	38.5	0.047	260	55	21
AF354558	321	42.4	0.003	81	25	30	AF386506	441	38.5	0.047	95	25	26
AY007306	1236	42.4	0.003	116	27	23	AF386505	442	38.5	0.047	95	25	26
AF015103	267	42.4	0.003	77	20	25	AE016747	300892	38.5	0.047	169	43	25
AL935252	343050	42	0.004	285	64	22	AE016747	300892	34.7	0.68	302	60	19
AL646086	92509	42	0.004	264	60	22	U25820	108	38.5	0.047	36	19	52
AL646086	92509	36.2	0.23	260	56	21	AF015111	257	38.5	0.047	74	22	29
AL646086	92509	35.8	0.31	247	57	23	AE005766	10091	38.1	0.062	73	21	28
AJ311994	206	41.6	0.006	65	24	36	Z31376	4192	38.1	0.062	29	19	65
J01801	1094	41.6	0.006	20	20	100	X86999	185	37.7	0.08	43	16	37
AF015095	267	41.6	0.006	74	24	32	X86998	185	37.7	0.08	43	16	37
AF015090	267	41.6	0.006	74	24	32	X86997	185	37.7	0.08	43	16	37
M58145	224	41.6	0.006	20	20	100	X86996	185	37.7	0.08	43	16	37
J01800	349	41.6	0.006	20	20	100	X86995	185	37.7	0.08	43	16	37
BA0000182814816	41.2	0.007	332	72	21	X86994	185	37.7	0.08	43	16	37	
AF015115	267	41.2	0.007	74	23	31	X86993	185	37.7	0.08	43	16	37
AF015113	257	41.2	0.007	74	23	31	X86992	185	37.7	0.08	43	16	37
AF015112	257	41.2	0.007	74	23	31	X86990	185	37.7	0.08	43	16	37
AF015102	257	41.2	0.007	74	23	31	X86989	185	37.7	0.08	43	16	37
AF015100	257	41.2	0.007	74	23	31	X86988	185	37.7	0.08	43	16	37
AF015099	257	41.2	0.007	74	23	31	X86987	185	37.7	0.08	43	16	37
AF015098	257	41.2	0.007	74	23	31	X86986	185	37.7	0.08	43	16	37
AF015096	257	41.2	0.007	74	23	31	X86985	185	37.7	0.08	43	16	37
AF015094	257	41.2	0.007	74	23	31	X86984	185	37.7	0.08	43	16	37
AF015091	257	41.2	0.007	74	23	31	X86983	185	37.7	0.08	43	16	37
AP003362	346300	41.2	0.007	332	72	21	X86982	185	37.7	0.08	43	16	37
Z54217	2262	40.8	0.009	20	20	100	X86981	185	37.7	0.08	43	16	37
AY575004	135	40.8	0.009	38	18	47	X86980	185	37.7	0.08	43	16	37
AY575002	136	40.8	0.009	38	18	47	X86979	185	37.7	0.08	43	16	37
AY575001	136	40.8	0.009	38	18	47	X87005	185	37.7	0.08	43	16	37
AF270112	3067	40.8	0.009	262	58	22	X87004	185	37.7	0.08	43	16	37
AF269857	3319	40.8	0.009	262	58	22	X87003	185	37.7	0.08	43	16	37
AF269729	3867	40.8	0.009	262	58	22	X87002	185	37.7	0.08	43	16	37
S62783	193	40.4	0.012	32	20	62	X87001	185	37.7	0.08	43	16	37
S62780	191	40.4	0.012	32	20	62	X87000	185	37.7	0.08	43	16	37
S62779	191	40.4	0.012	32	20	52	AL596172	248050	37.7	0.08	292	64	21
S62776	191	40.4	0.012	32	20	62	AE004867	15356	36.6	0.18	201	49	24
S62775	192	40.4	0.012	32	20	62	X85991	186	36.6	0.18	43	15	34
S62773	200	40.4	0.012	32	20	62	AY603345	483	36.6	0.18	166	34	20
AF525505	11500	40.4	0.012	278	64	23	AY603344	483	36.6	0.18	166	34	20
AF525505	11500	39.3	0.028	321	76	23	AY603342	483	35.6	0.18	166	34	20
AY575003	136	40.4	0.012	38	18	47	AY278535	360	35.6	0.18	115	31	26
AF015114	267	40.4	0.012	64	21	32	AE016827	2314078	36.2	0.23	273	58	21
AF015110	267	40.4	0.012	64	21	32	AE016827	2314078	35.2	0.23	273	58	21
AF015107	267	40.4	0.012	64	21	32	AE016827	2314078	35.6	0.31	274	61	22
AF015093	267	40.4	0.012	64	21	32	AY603343	483	38.2	0.23	166	34	20
AF015092	267	40.4	0.012	64	21	32	AY458638	19060	36.2	0.23	146	39	26
CP0000241796226	40	0.016	302	64	21	AE016958	302070	35.8	0.31	163	40	21	
CP0000231796846	40	0.016	302	64	21	HA0000196413771	35.8	0.31	286	59	20		
J01607	351	40	0.016	20	18	90	M12293	181	35.8	0.31	17	17	100
U96166	19841	39.7	0.021	278	48	17	AF459093	6852	35.4	0.4	269	58	21
U96239	390	39.7	0.021	97	26	26	BX294135	340750	35.4	0.4	240	52	21
U96238	390	39.7	0.021	97	26	26	AY278547	377	35.4	0.4	129	35	27
U96237	390	39.7	0.021	97	26	26	AE016940	304230	35	0.52	150	44	28
U96236	390	39.7	0.021	97	26	26	AE001129	10845	35	0.52	77	18	23
U96235	390	39.7	0.021	97	26	26	AY278538	27297	35	0.52	239	49	20
AY374137	491	39.7	0.021	97	26	26	AF045472	7263	34.7	0.68	95	25	26
AY374135	485	39.7	0.021	97	26	26	AE001578	29838	34.7	0.68	114	30	26
AY374134	485	39.7	0.021	97	26	26	BA0000163031430	34.7	0.68	232	49	21	
AY374133	485	39.7	0.021	97	26	26	BA0000454659019	34.7	0.68	73	24	32	
AY374132	485	39.7	0.021	97	26	26	U40259	8013	34.7	0.68	95	25	26
EX5718572799802	39.3	0.028	283	66	23	AE016864	310325	34.3	0.89	128	31	24	
BA0000332820462	39.3	0.028	283	66	23	AE014627	10029	34.3	0.89	271	54	19	

Fig. 25A-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AE017201	300478	33.9	1.2	125	28	22							
AE011395	10368	33.9	1.2	59	18	30							
AE015945	89370	33.9	1.2	103	24	23							
AL935255	269050	33.9	1.2	299	58	19							
CR378673	349814	33.9	1.2	73	16	21							
AE017310	300343	33.9	1.2	171	36	21							
AB015410	3519	33.9	1.2	283	55	19							
AE016858	312639	33.1	2	162	41	25							
AE016813	336182	33.1	2	120	30	25							
gi 42627766 3786		33.1	2	62	19	30							
AE016748	300029	33.1	2	148	30	20							
CR378667	349080	33.1	2	137	31	22							
AF113610	3246	33.1	2	240	48	20							
AF322013	230573	33.1	2	62	19	30							
D90775	16902	33.1	2	147	37	26							
D90774	18700	33.1	2	147	37	25							
CR378676	343629	33.1	2	107	27	25							
AE017135	291326	32.7	2.6	290	61	21							
AY708386	1203	32.7	2.6	254	52	20							
AJ414152	313050	32.7	2.6	290	61	21							
AE013773	10520	32.7	2.6	290	61	21							
AE005723	10573	32.7	2.6	235	47	20							
AE017240	299986	32.7	2.6	70	21	30							

Fig. 25B

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
CAE53942	349	386	1.00E-106	348	215	61	AAA17862	505	336	2.00E-91	178	177	99
CAD60547	349	385	1.00E-106	348	214	61	AAA17862	505	197	7.00E-50	227	126	55
AAP13297	349	384	1.00E-106	348	213	61	AAA17859	508	336	2.00E-91	178	177	99
CAE53943	349	383	1.00E-105	348	214	61	AAA17859	508	197	1.00E-49	305	140	45
CAD56695	349	382	1.00E-105	348	213	61	AAR10747	505	335	4.00E-91	178	176	98
AAU90046	271	379	1.00E-104	284	202	71	AAR10747	505	194	6.00E-49	227	125	55
CAA62509	313	369	1.00E-101	313	206	65	AAA17865	508	335	4.00E-91	178	176	98
BAA06987	348	367	1.00E-101	356	214	60	AAA17865	508	193	2.00E-48	106	103	97
NP_929221	355	364	1.00E-100	355	207	58	AAR10749	504	334	5.00E-91	178	176	98
AAA62397	357	356	1.00E-97	367	213	58	AAR10749	504	189	3.00E-47	106	101	95
AAA62396	355	355	2.00E-97	365	211	57	AAR10748	504	334	5.00E-91	178	176	98
ZP_00169997361	343	1.00E-93	361	201	55	AAR10748	504	190	1.00E-47	280	139	49	
AAT81649	505	336	2.00E-91	178	177	99	AAR10594	505	334	5.00E-91	178	176	98
AAT81649	505	194	8.00E-49	106	104	98	AAR10594	505	194	6.00E-49	227	125	55
AAT81648	505	336	2.00E-91	178	177	99	AAT81624	504	334	7.00E-91	178	176	98
AAT81648	505	194	8.00E-49	106	104	98	AAT81624	504	191	9.00E-48	301	136	45
AAT81622	505	336	2.00E-91	178	177	99	AAT81623	504	334	7.00E-91	178	176	98
AAT81622	505	197	7.00E-50	227	126	55	AAT81623	504	191	9.00E-48	301	136	45
AAT81621	505	336	2.00E-91	178	177	99	CAA78794	504	334	7.00E-91	177	176	99
AAT81621	505	195	4.00E-49	227	125	55	CAA78794	504	196	3.00E-49	227	126	55
AAT81620	505	336	2.00E-91	178	177	99	CAA78781	504	334	7.00E-91	177	176	99
AAT81620	505	196	3.00E-49	227	126	55	CAA78781	504	197	7.00E-50	227	125	55
AAT81619	505	336	2.00E-91	178	177	99	CAA78780	504	334	7.00E-91	177	176	99
AAT81619	505	194	8.00E-49	105	104	98	CAA78780	504	194	8.00E-49	106	104	98
AAT81618	505	336	2.00E-91	178	177	99	CAA78779	507	334	7.00E-91	177	176	99
AAT81618	505	194	8.00E-49	106	104	98	CAA78779	507	194	8.00E-49	106	104	98
AAT81617	505	336	2.00E-91	178	177	99	CAA78778	504	334	7.00E-91	177	176	99
AAT81617	505	194	8.00E-49	106	104	98	CAA78778	504	194	8.00E-49	106	104	98
AAT81614	505	336	2.00E-91	178	177	99	CAA78777	504	334	7.00E-91	177	176	99
AAT81614	505	197	1.00E-49	227	126	55	CAA78777	504	194	8.00E-49	106	104	98
AAT81613	505	336	2.00E-91	178	177	99	CAA78776	504	334	7.00E-91	177	176	99
AAT81613	505	194	6.00E-49	227	125	55	CAA78776	504	194	8.00E-49	106	104	98
AAT81612	505	336	2.00E-91	178	177	99	CAA78775	504	334	7.00E-91	177	176	99
AAT81612	505	197	1.00E-49	227	126	55	CAA78775	504	197	1.00E-49	227	126	55
AAR10745	508	336	2.00E-91	178	177	99	CAA78774	504	334	7.00E-91	177	176	99
AAR10745	508	197	7.00E-50	324	142	43	CAA78774	504	197	7.00E-50	227	126	55
AAR10628	504	336	2.00E-91	178	177	99	CAA78773	507	334	7.00E-91	177	176	99
AAR10628	504	194	6.00E-49	309	141	45	CAA78773	507	197	7.00E-50	324	142	43
AAR10624	508	336	2.00E-91	178	177	99	S33187	504	334	7.00E-91	177	176	99
AAR10624	508	194	8.00E-49	106	104	98	S33187	504	197	1.00E-49	227	126	55
AAR10622	508	336	2.00E-91	178	177	99	AAA53497	507	334	7.00E-91	177	176	99
AAR10622	508	194	8.00E-49	106	104	98	AAA53497	507	194	8.00E-49	106	104	98
AAR10621	508	336	2.00E-91	178	177	99	AAA53495	504	334	7.00E-91	177	176	99
AAR10621	508	194	8.00E-49	106	104	98	AAA53495	504	194	8.00E-49	106	104	98
AAR10619	505	336	2.00E-91	178	177	99	AAA53491	507	334	7.00E-91	177	176	99
AAR10619	505	197	7.00E-50	227	126	55	AAA53491	507	194	8.00E-49	106	104	98
AAR10618	505	335	2.00E-91	178	177	99	AAA53490	504	334	7.00E-91	177	176	99
AAR10618	505	197	7.00E-50	227	126	55	AAA53490	504	196	2.00E-49	227	126	55
AAR10616	505	335	2.00E-91	178	177	99	AAT81603	505	333	9.00E-91	178	176	98
AAR10616	505	194	8.00E-49	106	104	98	AAT81603	505	194	8.00E-49	106	104	98
AAR10610	505	335	2.00E-91	178	177	99	AAA27088	505	333	9.00E-91	178	176	98
AAR10610	505	194	8.00E-49	106	104	98	AAA27088	505	192	2.00E-48	106	103	97
AAR10609	504	336	2.00E-91	178	177	99	JUC056	351	333	1.00E-90	354	196	55
AAR10609	504	193	2.00E-48	106	103	97	AAA53493	465	330	7.00E-90	177	175	98
AAR10608	504	336	2.00E-91	178	177	99	AAA53493	465	171	1.00E-41	91	91	100
AAR10608	504	193	2.00E-48	106	103	97	AAR10625	504	330	1.00E-89	178	173	97
AAR10604	505	336	2.00E-91	178	177	99	AAR10625	504	187	1.00E-46	227	120	52
AAR10604	505	194	8.00E-49	106	104	98	AAA53496	503	328	4.00E-89	177	172	97
O52959	505	336	2.00E-91	178	177	99	AAA53496	503	187	1.00E-46	227	120	52
O62959	505	194	8.00E-49	106	104	98	YP_049831	290	327	8.00E-89	302	187	61
AAA27092	505	336	2.00E-91	178	177	99	S78461	364	319	2.00E-86	365	203	55
AAA27092	505	194	8.00E-49	106	104	98	NP_992909	369	318	5.00E-86	368	202	54
AAA27090	505	336	2.00E-91	178	177	99	NP_405406	369	318	5.00E-86	368	202	54
AAA27090	505	194	8.00E-49	106	104	98	AOB86383	369	315	2.00E-85	368	201	54
AAA27085	494	336	2.00E-91	178	177	99	AAA64387	493	315	4.00E-85	165	165	99
AAA27085	494	174	9.00E-43	95	93	97	AAA64387	493	194	8.00E-49	106	104	98
AAA17863	508	336	2.00E-91	178	177	99	AAQ22673	488	311	4.00E-84	263	178	67
AAA17863	508	194	6.00E-49	227	125	55	AAQ22673	488	160	1.00E-38	103	83	80

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AAR10689	505	311	4.00E-84	244	175	71	A48658	595	135	5.00E-31	113	70	61
AAR10689	505	175	5.00E-43	140	96	68	AAP13337	436	296	2.00E-79	300	175	58
BAD16577	488	310	8.00E-84	273	180	65	AAP13337	436	135	6.00E-31	92	67	72
BAD16577	488	165	3.00E-40	280	121	43	AAL30164	498	296	2.00E-79	178	155	87
S78460	358	310	1.00E-83	359	194	54	AAL30164	498	155	3.00E-37	174	94	54
AAQ22680	488	309	2.00E-83	263	177	67	AAR97969	378	296	2.00E-79	223	159	71
AAQ22680	488	160	1.00E-38	103	83	80	BAA85089	565	296	2.00E-79	302	176	58
AAL30165	487	308	4.00E-83	256	177	69	BAA85089	565	133	2.00E-30	102	66	64
AAL30165	487	158	7.00E-38	103	82	79	AAP13309	554	296	2.00E-79	237	161	67
CAD99230	454	308	5.00E-83	262	173	66	AAP13309	554	141	8.00E-33	295	105	35
CAD99230	454	117	1.00E-25	75	59	78	AAF85764	554	296	2.00E-79	237	161	67
AAR10691	505	308	5.00E-83	244	172	70	AAF85764	554	139	3.00E-32	293	104	35
AAR10691	505	171	7.00E-42	140	94	67	AAQ22687	503	295	3.00E-79	178	155	87
AAR10690	505	308	5.00E-83	244	172	70	AAQ22687	503	158	7.00E-38	265	111	41
AAR10690	505	173	2.00E-42	140	95	67	AAP13324	545	295	3.00E-79	218	157	72
AAQ22683	525	306	1.00E-82	299	184	61	AAP13324	545	140	1.00E-32	172	85	49
AAQ22683	525	149	4.00E-35	92	76	82	AAP13324	545	32	7.10E+00	219	51	23
BAD14980	524	306	1.00E-82	299	184	61	CAA85351	565	285	3.00E-79	222	156	70
BAD14980	524	150	1.00E-35	92	77	83	CAA85351	565	133	2.00E-30	102	66	64
AAQ22684	525	305	3.00E-82	299	184	61	AAQ22689	557	295	3.00E-79	284	169	59
AAQ22684	525	149	4.00E-35	92	76	82	AAQ22689	557	141	8.00E-33	115	71	61
AAB17947	585	305	3.00E-82	274	170	62	AAP13336	447	295	3.00E-79	274	171	62
AAB17947	585	142	5.00E-33	115	74	64	AAP13336	447	134	1.00E-30	92	67	72
S44982	524	305	4.00E-82	299	183	61	AAP13318	579	295	3.00E-79	236	161	68
S44982	524	148	5.00E-35	92	76	82	AAP13318	579	140	1.00E-32	116	72	62
AAG56938	585	305	4.00E-82	274	169	61	AAP13316	557	295	3.00E-79	264	169	59
AAG56938	585	142	5.00E-33	115	74	64	AAP13316	557	142	5.00E-33	115	71	61
AAQ22676	585	305	4.00E-82	274	169	61	AAP13307	662	295	3.00E-79	257	168	65
AAQ22676	585	142	5.00E-33	115	74	64	AAP13307	662	142	3.00E-33	273	102	37
AAQ22676	585	305	4.00E-82	267	168	62	AAQ13298	436	295	4.00E-79	291	174	59
AAQ22676	585	142	5.00E-33	115	74	64	AAQ13298	436	135	6.00E-31	92	67	72
AAF71887	585	305	4.00E-82	267	168	62	AAL30167	545	295	4.00E-79	218	157	72
AAF71887	585	142	5.00E-33	115	74	64	AAL30167	545	140	1.00E-32	172	85	49
AAQ22685	525	304	6.00E-82	299	183	61	AAQ22681	447	294	6.00E-79	290	171	58
AAQ22685	525	149	4.00E-35	92	76	82	AAQ22681	447	137	1.00E-31	314	115	36
AAQ22685	525	304	7.00E-82	274	169	61	AAQ13226	447	294	6.00E-79	290	171	58
AAF71901	585	305	5.00E-33	115	74	64	AAQ13226	447	137	2.00E-31	314	114	36
AAF71901	585	142	5.00E-33	115	74	64	AAP13302	670	284	6.00E-79	247	162	65
C48658	584	303	2.00E-81	267	169	63	AAP13302	670	147	2.00E-34	312	104	33
C48658	584	141	8.00E-33	115	73	63	AAP13334	564	294	8.00E-79	222	156	70
BAD14957	423	301	6.00E-81	296	186	62	AAP13334	564	133	2.00E-30	102	66	64
BAD14957	423	141	8.00E-33	87	75	88	AAP13334	564	294	8.00E-79	222	156	70
AAQ13299	548	300	1.00E-80	266	171	64	AAL30168	564	294	8.00E-79	222	156	70
AAQ13299	548	140	1.00E-32	102	69	67	AAL30168	564	132	4.00E-30	102	66	64
AAQ13321	555	299	2.00E-80	295	171	57	NP_707809	550	293	1.00E-78	202	154	76
AAQ13321	555	140	1.00E-32	116	71	61	NP_707809	550	140	1.00E-32	116	71	61
B48588	595	298	3.00E-80	223	157	70	AAQ22690	440	293	1.00E-78	288	177	61
B48588	595	139	3.00E-32	255	106	41	AAQ22690	440	137	2.00E-31	312	109	34
AAQ22682	595	298	3.00E-80	223	157	70	AAN34779	498	293	1.00E-78	178	153	85
AAQ22682	595	137	1.00E-31	113	71	62	AAN34779	498	155	3.00E-37	174	94	54
AAQ22679	595	298	3.00E-80	223	157	70	AAP13311	576	293	1.00E-78	299	171	57
AAQ22679	595	137	1.00E-31	113	71	62	AAP13311	576	142	4.00E-33	287	111	38
NP_754230	595	298	3.00E-80	223	157	70	AAP13332	572	293	1.00E-78	229	160	69
NP_754230	595	138	5.00E-32	255	106	41	AAP13332	572	144	1.00E-33	116	73	62
AAQ13331	568	298	3.00E-80	223	160	71	AAP13332	572	32	7.10E+00	221	44	19
AAQ13331	568	140	1.00E-32	114	74	64	AAP13306	550	293	1.00E-78	202	154	76
AAQ13305	595	298	3.00E-80	223	157	70	AAP13306	550	140	1.00E-32	116	71	61
AAQ13305	595	137	1.00E-31	113	71	62	AAF32260	550	293	1.00E-78	202	154	76
BAA85086	568	298	3.00E-80	223	160	71	AAF32260	550	138	7.00E-32	276	102	36
BAA85086	568	141	6.00E-33	270	104	38	AAQ22677	436	293	2.00E-78	291	169	58
CAD97427	545	298	5.00E-80	246	163	66	AAQ22677	436	134	8E-31	92	67	72
CAD97427	545	91.3	1.00E-17	84	45	53	AAP13328	559	292	2.00E-78	179	149	83
AAP13313	588	297	9.00E-80	219	162	73	AAP13328	559	139	4.00E-32	113	69	61
AAP13313	588	142	3.00E-33	149	83	55	AAP13320	443	291	4.00E-78	288	176	61
AAQ22674	447	298	1.00E-79	290	172	59	AAP13320	443	136	5.00E-31	315	107	33
AAQ22674	447	135	3.00E-31	181	87	48	AAN34780	498	291	5.00E-78	178	152	85
AAP13335	605	296	1.00E-79	222	157	70	AAN34780	498	155	3.00E-37	174	94	54
AAP13335	605	139	3.00E-32	163	81	49	AAR10708	413	291	5.00E-78	275	175	63
A48658	595	296	2.00E-79	223	156	69	AAR10708	413	150	1E-35	100	79	79

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AAR10698	421	291	5.00E-78	276	170	61	AAR10715	426	149	2.00E-35	174	91	52
AAR10698	421	152	5.00E-36	102	79	77	AAR10711	422	288	5.00E-77	214	161	75
BAA05151	568	291	5.00E-78	263	159	60	AAR10711	422	150	1.00E-35	100	79	79
BAA05151	568	140	1.00E-32	114	74	64	AAR10706	422	268	5.00E-77	214	161	75
AAQ22686	610	291	6.00E-78	191	151	79	AAR10706	422	150	1.00E-35	100	79	79
AAQ22686	610	145	4.00E-34	287	109	37	AAR10666	501	288	5.00E-77	216	158	73
AAP13322	570	291	6.00E-78	179	150	83	AAR10666	501	156	2.00E-37	100	82	82
AAP13322	570	139	3.00E-32	234	88	37	AAR10659	500	288	5.00E-77	208	157	75
AAP13310	613	291	6.00E-78	191	151	79	AAR10659	500	156	2.00E-37	100	82	82
AAP13310	613	146	3.00E-34	285	109	38	AAR10658	500	288	5.00E-77	208	157	75
CAD99229	453	291	6.00E-78	251	164	65	AAR10558	500	156	2.00E-37	100	82	82
CAD99229	453	117	1.00E-25	75	59	78	AAR10543	500	288	5.00E-77	208	157	75
BAA85087	570	290	1.00E-77	229	159	69	AAR10543	500	154	7.00E-37	100	81	81
BAA85087	570	144	1.00E-33	116	73	62	AAR10667	501	287	7.00E-77	216	158	73
BAA85087	570	32	7.10E+00	221	44	19	AAR10667	501	156	2.00E-37	100	82	82
BAD14961	529	290	1.00E-77	295	177	59	NP_416433	498	287	9.00E-77	177	146	82
BAD14961	529	151	8.00E-36	293	112	38	NP_416433	498	149	4.00E-35	151	80	52
AAP13327	432	290	1.00E-77	286	169	59	AAQ22678	426	287	9.00E-77	258	163	63
AAP13327	432	136	2.00E-31	253	102	40	AAQ22678	426	138	7.00E-32	232	93	40
AAP13323	428	290	1.00E-77	287	166	62	AAT81632	500	287	9.00E-77	206	157	76
AAP13323	428	134	1.00E-30	127	74	58	AAT81632	500	152	4.00E-36	98	79	80
AAL30166	428	290	1.00E-77	287	166	62	AAT81631	500	287	9.00E-77	198	154	77
AAL30166	428	131	7.00E-30	127	73	57	AAT81631	500	156	2.00E-37	100	82	82
AAR10700	421	290	1.00E-77	271	170	62	CAA35488	498	287	9.00E-77	177	146	82
AAR10700	421	152	5.00E-36	102	79	77	CAA35488	498	151	6.00E-36	244	101	41
AAR10699	421	290	1.00E-77	271	170	62	AAR10755	422	287	9.00E-77	211	159	75
AAR10699	421	152	5.00E-36	102	79	77	AAR10755	422	150	1.00E-35	100	79	79
AAR10695	421	290	1.00E-77	175	153	87	AAR10669	493	287	9.00E-77	198	154	77
AAR10695	421	289	2.00E-77	175	153	87	AAR10669	493	156	2.00E-37	100	82	82
AAR10696	421	152	5.00E-36	102	79	77	AAR10668	481	287	9.00E-77	245	162	66
AAR10694	421	289	2.00E-77	175	153	87	AAR10568	481	156	2.00E-37	100	82	82
AAP13325	561	289	2.00E-77	233	162	69	AAR10565	501	287	9.00E-77	206	157	76
AAP13325	561	150	1.00E-35	279	102	36	AAR10665	501	156	2.00E-37	100	82	82
AAR10759	422	289	2.00E-77	284	175	61	AAR10564	501	287	9.00E-77	206	157	76
AAR10759	422	150	1.00E-35	100	79	79	AAR10664	501	154	7.00E-37	100	81	81
AAR10544	500	288	3.00E-77	208	157	75	AAR10662	500	287	9.00E-77	206	157	76
AAR10544	500	155	4.00E-37	100	81	AAR10662	500	156	2.00E-37	100	82	82	
BAA05153	549	288	3.00E-77	234	159	67	AAR10660	500	287	9.00E-77	198	154	77
BAA05153	549	135	6.00E-31	240	86	35	AAR10660	500	156	2.00E-37	100	82	82
AAP13329	426	288	4.00E-77	258	164	63	AAR10590	502	287	9.00E-77	220	160	72
AAP13329	426	138	7.00E-32	232	93	40	AAR10590	502	157	1.00E-37	100	82	82
AAP13319	570	288	4.00E-77	179	149	83	AAR10721	421	286	1.00E-76	175	152	86
AAP13319	570	139	4.00E-32	237	91	38	AAR10721	421	150	1.00E-35	100	79	79
AAF32251	426	288	4.00E-77	258	164	63	AAR10713	421	286	1.00E-76	175	152	86
AAF32281	426	136	3.00E-31	92	69	75	AAR10713	421	150	1.00E-35	100	79	79
AAR10710	422	288	4.00E-77	237	164	69	AAR10557	500	286	1.00E-76	198	154	77
AAR10710	422	148	5.00E-35	100	78	78	AAR10557	500	152	4.00E-36	98	79	80
AAT81644	426	288	5.00E-77	214	161	75	AAR10572	421	286	1.00E-76	175	152	86
AAT81644	426	150	1.00E-35	100	79	79	AAR10721	421	150	1.00E-35	100	79	79
AAT81643	422	288	5.00E-77	214	161	75	AAR10713	421	286	1.00E-76	175	152	86
AAT81643	422	148	5.00E-35	100	78	78	AAR10713	421	150	1.00E-35	100	79	79
AAT81642	424	288	5.00E-77	214	161	75	AAR10557	500	286	1.00E-76	198	154	77
AAT81642	424	150	1.00E-35	100	79	79	AAR10557	500	152	4.00E-36	98	79	80
AAT81611	502	288	5.00E-77	220	160	72	AAT81609	502	286	2.00E-76	220	159	72
AAT81611	502	157	1.00E-37	100	82	82	AAT81609	502	157	1.00E-37	100	82	82
AAT81610	502	288	5.00E-77	220	160	72	AAR10562	499	286	2.00E-76	206	157	76
AAT81610	502	157	1.00E-37	100	82	82	AAR10562	499	158	5.00E-38	338	127	37
AAP13330	501	288	5.00E-77	220	156	70	AAR10561	499	286	2.00E-76	209	158	75
AAP13330	501	134	1.00E-30	151	77	50	AAR10561	499	158	7.00E-38	338	127	37
AAR10761	422	288	5.00E-77	214	161	75	AAR10559	499	286	2.00E-76	209	158	75
AAR10761	422	148	5.00E-35	100	78	78	AAR10559	499	158	7.00E-38	338	127	37
AAR10760	422	288	5.00E-77	214	161	75	AAR10558	499	157	9.00E-38	338	127	37
AAR10760	422	150	1.00E-35	100	79	79	AAR10557	499	286	2.00E-76	209	158	75
AAR10719	422	288	5.00E-77	214	161	75	AAR10557	499	158	7.00E-38	338	127	37
AAR10719	422	150	1.00E-35	100	79	79	AAR10557	499	286	2.00E-76	209	158	75
AAR10717	426	288	5.00E-77	214	161	75	AAF32259	420	285	4.00E-76	269	160	59
AAR10717	426	151	8.00E-36	174	92	52	AAF32259	420	135	6.00E-31	138	77	55
AAR10715	426	288	5.00E-77	214	161	75	AAR10714	421	285	4.00E-76	175	151	86

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AAR10714	421	150	1.00E-35	100	79	79	AAR10739	506	154	7.00E-37	100	80	80
AAR10572	495	285	4.00E-76	306	176	57	AAR10512	506	282	3.00E-75	232	159	68
AAR10572	495	149	3.00E-35	89	76	85	AAR10512	506	155	4.00E-37	100	81	81
NP_841632	275	285	5.00E-76	297	159	53	AAR10511	506	282	3.00E-75	232	159	68
CAD97429	408	285	5.00E-76	179	145	81	AAR10511	506	155	4.00E-37	100	81	81
CAD97429	408	85.7	2.00E-16	285	82	28	AAR10489	506	282	3.00E-75	232	159	68
AAT81605	495	285	5.00E-76	306	176	57	AAR10489	506	155	4.00E-37	100	81	81
AAT81605	495	151	6.00E-36	96	78	81	AAR10488	506	282	3.00E-75	232	159	68
AAP13303	420	285	5.00E-76	189	147	77	AAR10488	506	155	4.00E-37	100	81	81
AAP13303	420	137	2.00E-31	138	78	56	AAC43352	506	281	4.00E-75	228	156	68
AAR10720	422	285	5.00E-76	211	158	74	AAC43352	506	155	4.00E-37	100	81	81
AAR10720	422	150	1.00E-35	100	79	79	AAR10504	506	281	4.00E-75	268	163	60
AAR10575	495	285	5.00E-76	306	175	57	AAR10504	506	155	4.00E-37	100	81	81
AAR10575	495	151	6.00E-36	96	78	81	AAR10495	506	281	4.00E-75	268	162	60
AAR10574	495	285	5.00E-76	306	176	57	AAR10495	506	155	4.00E-37	100	81	81
AAR10574	495	151	6.00E-36	96	78	81	AAR10754	494	281	5.00E-75	175	148	84
AAR10515	501	285	5.00E-76	175	150	85	AAR10754	494	152	4.00E-36	98	79	80
AAR10515	501	155	4.00E-37	100	81	81	AAC43351	506	281	5.00E-75	175	148	84
AAR10729	501	284	6.00E-76	215	157	73	AAC43351	506	155	4.00E-37	100	81	81
AAR10729	501	152	4.00E-36	98	79	80	AAC43348	506	281	5.00E-76	175	148	84
AAR10728	501	284	6.00E-76	215	157	73	AAC43348	506	154	7.00E-37	100	81	81
AAR10728	501	152	4.00E-36	98	79	80	AAR10751	495	281	5.00E-75	175	148	84
AAR10716	426	284	6.00E-76	214	160	74	AAR10751	495	148	7.00E-35	98	77	78
AAR10716	426	149	2.00E-35	174	91	52	AAR10750	495	281	5.00E-75	175	148	84
AAR10702	499	284	6.00E-76	192	152	79	AAR10750	495	152	4.00E-36	98	79	80
AAR10702	499	153	2.00E-36	100	80	80	AAR10743	506	281	5.00E-75	175	148	84
CAA27130	509	284	8.00E-76	175	150	85	AAR10743	506	155	4.00E-37	100	81	81
CAA27130	509	149	4.00E-35	91	76	83	AAR10742	506	281	5.00E-75	175	148	84
NP_456520	506	284	8.00E-76	175	150	85	AAR10742	506	155	4.00E-37	100	81	81
NP_456520	506	149	4.00E-35	91	76	83	AAR10741	506	281	5.00E-75	175	148	84
AAQ22688	420	284	8.00E-76	189	147	77	AAR10741	506	154	1.00E-36	100	80	80
AAQ22688	420	135	6.00E-31	138	77	55	AAR10740	506	281	5.00E-75	175	148	84
AAT81639	506	284	8.00E-76	175	150	85	AAR10740	506	155	6.00E-37	100	81	81
AAT81639	506	149	4.00E-35	91	76	83	AAR10730	495	281	5.00E-75	175	148	84
AAT81608	506	284	8.00E-76	175	150	85	AAR10730	495	155	6.00E-37	245	105	42
AAT81608	506	149	4.00E-35	91	76	83	AAR10724	494	281	5.00E-75	175	148	84
AAT81607	506	284	8.00E-76	175	150	85	AAR10724	494	158	5.00E-38	272	108	39
AAT81607	506	150	1.00E-35	91	77	84	AAR10682	499	281	5.00E-75	175	148	84
AAR10646	419	284	8.00E-76	175	150	85	AAR10682	499	154	1.00E-36	100	80	80
AAR10646	419	149	4.00E-35	91	76	83	AAR10681	499	281	5.00E-75	175	148	84
AAR10588	506	284	8.00E-76	175	150	85	AAR10681	499	154	1.00E-36	100	80	80
AAR10588	506	150	1.00E-35	91	77	84	AAR10648	495	281	5.00E-76	175	148	84
AAR10514	506	284	8.00E-76	175	150	85	AAR10648	495	152	4.00E-36	98	79	80
AAR10514	506	150	1.00E-35	91	77	84	AAR10567	506	281	5.00E-75	175	148	84
S09638	505	284	8.00E-76	175	150	85	AAR10567	506	155	4.00E-37	100	81	81
S09638	505	149	4.00E-35	91	76	83	AAR10566	500	281	5.00E-75	175	148	84
NP_461698	506	283	1.00E-75	232	160	68	AAR10566	500	157	1.00E-37	100	82	62
NP_461698	506	155	4.00E-37	100	81	81	AAR10564	500	281	5.00E-75	175	148	84
YP_151805	506	283	1.00E-75	279	165	59	AAR10564	500	155	4.00E-37	100	81	81
YP_151805	506	155	4.00E-37	100	81	81	AAR10542	495	281	5.00E-75	175	148	84
AAR10471	501	283	1.00E-75	295	169	57	AAR10542	495	155	6.00E-37	100	81	81
AAR10471	501	151	6.00E-36	98	78	79	AAR10539	501	281	5.00E-75	175	148	84
AAR10484	506	283	1.00E-75	232	160	68	AAR10539	501	155	4.00E-37	100	81	81
AAR10484	506	155	4.00E-37	100	81	81	AAR10538	501	281	5.00E-75	175	148	84
CAB65960	456	283	2.00E-75	247	162	65	AAR10538	501	155	4.00E-37	100	81	81
CAB65960	456	139	3.00E-32	88	71	80	AAR10536	501	281	5.00E-75	175	148	84
AAR10737	506	283	2.00E-75	229	160	69	AAR10536	501	155	4.00E-37	100	81	81
AAR10737	506	154	9.00E-37	150	87	58	AAR10532	501	281	5.00E-75	175	148	84
AAR10735	506	283	2.00E-75	229	160	69	AAR10532	501	155	4.00E-37	100	81	81
AAR10735	506	154	1.00E-36	100	80	80	AAR10530	501	281	5.00E-75	175	148	84
AAR10703	499	282	2.00E-75	199	154	77	AAR10530	501	155	4.00E-37	100	81	81
AAR10703	499	151	8.00E-36	98	78	79	AAR10527	501	281	5.00E-75	175	148	84
AAR10499	506	282	2.00E-75	275	164	59	AAR10527	501	155	4.00E-37	100	81	81
AAR10499	506	155	4.00E-37	100	81	81	AAR10523	501	281	5.00E-75	175	148	84
AAR10509	505	282	3.00E-75	232	159	68	AAR10523	501	155	4.00E-37	100	81	81
AAR10509	505	155	4.00E-37	100	81	81	AAR10521	501	281	5.00E-75	175	148	84
CAD22870	280	282	3.00E-75	296	167	56	AAR10521	501	155	6E-37	100	81	81
AAR10739	506	282	3.00E-75	217	156	71	AAR10518	501	281	5.00E-75	175	148	84

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AAR10518	501	155	6.00E-37	100	81	81	AAR10684	499	154	9.00E-37	98	80	81
AAR10505	465	281	5.00E-75	175	148	84	AAR10583	499	280	9.00E-75	175	148	84
AAR10505	465	155	4.00E-37	100	81	81	AAR10683	499	154	9.00E-37	98	80	81
AAR10501	506	281	5.00E-75	175	148	84	AAR10679	499	280	9.00E-75	175	148	84
AAR10501	506	155	4.00E-37	100	81	81	AAR10679	499	155	4.00E-37	100	81	81
AAR10496	506	281	5.00E-75	175	148	84	AAR10578	506	159	4.00E-38	100	83	83
AAR10494	506	155	4.00E-37	100	81	81	AAR10677	506	280	9.00E-75	175	148	84
AAR10494	506	281	5.00E-75	175	148	84	AAR10677	506	158	5.00E-38	98	82	83
AAR10674	493	281	7.00E-75	229	160	69	AAR10676	491	280	9.00E-75	175	148	84
AAR10674	493	147	1.00E-34	98	77	78	AAR10676	491	152	4.00E-36	98	79	80
AAR10673	493	281	7.00E-75	229	160	69	AAR10675	491	280	9.00E-75	175	148	84
AAR10673	493	152	4.00E-36	98	79	80	AAR10675	491	152	4.00E-36	98	79	80
AAR10551	499	281	7.00E-75	217	157	72	AAR10653	495	280	9.00E-75	175	148	84
AAR10551	499	155	6.00E-37	100	81	81	AAR10653	495	157	1.00E-37	245	106	43
AAR10548	499	281	7.00E-75	217	157	72	AAR10644	495	280	9.00E-75	175	148	84
AAR10548	499	155	6.00E-37	100	81	81	AAR10644	495	152	4.00E-36	98	79	80
CAA27129	493	280	9.00E-75	175	148	84	AAR10637	495	280	9.00E-75	175	148	84
CAA27129	493	154	1.00E-36	329	126	38	AAR10637	495	152	4.00E-36	98	79	80
CAA27128	498	280	9.00E-75	175	148	84	AAR10633	495	280	9.00E-75	175	148	84
CAA27128	498	157	9.00E-38	100	83	83	AAR10633	495	152	4.00E-36	98	79	80
AAR10534	501	280	9.00E-75	175	148	84	AAR10581	501	280	9.00E-75	175	148	84
AAR10534	501	155	4.00E-37	100	81	81	AAR10581	501	152	4.00E-36	98	79	80
AAR10503	506	280	9.00E-75	175	148	84	AAR10579	501	280	9.00E-75	175	148	84
AAR10503	506	154	7.00E-37	98	80	81	AAR10579	501	154	7.00E-37	100	81	81
NP_460912	495	280	9.00E-75	175	148	84	AAR10576	501	280	9.00E-75	175	148	84
NP_460912	495	152	4.00E-36	98	79	80	AAR10576	501	154	7.00E-37	100	81	81
AAT81647	501	280	9.00E-75	175	148	84	AAR10569	495	280	9.00E-75	175	148	84
AAT81647	501	154	7.00E-37	100	81	81	AAR10569	495	158	2.00E-37	100	82	82
AAT81646	499	280	9.00E-75	175	148	84	AAR10568	495	280	9.00E-75	175	148	84
AAT81646	499	155	4.00E-37	100	81	81	AAR10568	495	156	2.00E-37	100	82	82
AAT81641	499	280	9.00E-75	175	148	84	AAR10563	498	280	9.00E-75	175	148	84
AAT81641	499	154	9.00E-37	98	80	81	AAR10563	498	155	6.00E-37	100	81	81
AAT81640	495	280	9.00E-75	175	148	84	AAR10555	495	280	9.00E-75	175	148	84
AAT81640	495	152	4.00E-36	98	79	80	AAR10555	495	155	6.00E-37	100	81	81
AAT81638	493	280	9.00E-75	175	148	84	AAR10553	495	280	9.00E-75	175	148	84
AAT81638	493	152	4.00E-36	98	79	80	AAR10553	495	155	4.00E-37	100	81	81
AAT81629	495	280	9.00E-75	175	148	84	AAR10525	501	280	9.00E-75	175	148	84
AAT81629	495	158	2.00E-37	245	106	43	AAR10525	501	155	6.00E-37	100	81	81
AAT81627	495	280	9.00E-75	175	148	84	AAR10524	501	280	9.00E-75	175	148	84
AAT81627	495	152	4.00E-36	98	79	80	AAR10524	501	155	6.00E-37	100	81	81
AAT81605	501	280	9.00E-75	175	148	84	AAR10513	495	280	9.00E-75	175	148	84
AAT81606	501	154	7.00E-37	100	81	81	AAR10513	495	155	6.00E-37	100	81	81
AAT81604	495	280	9.00E-75	175	148	84	AAK20804	504	280	9.00E-75	173	148	85
AAT81604	495	156	2.00E-37	100	82	82	AAK20804	504	149	4.00E-35	91	78	83
AAT68767	491	280	9.00E-75	175	148	84	S07276	494	280	9.00E-75	175	148	84
AAT68767	491	136	2.00E-31	90	71	78	S07276	494	157	9.00E-38	100	83	83
CAA28190	493	280	9.00E-75	175	148	84	S09637	489	280	9.00E-75	175	148	84
CAA28190	493	152	4.00E-36	98	79	80	S09637	489	154	1.00E-36	329	126	38
AAO13791	501	280	9.00E-75	175	148	84	AAF80752	501	280	1.00E-74	175	148	84
AAO13791	501	154	1.00E-36	100	80	80	AAF80752	501	154	7.00E-37	100	81	81
AAL18053	487	280	9.00E-75	175	148	84	AAR10652	495	280	1.00E-74	175	147	84
AAL18053	487	129	4.00E-29	85	66	77	AAR10652	495	152	4.00E-36	98	79	80
AAR10738	506	280	9.00E-75	175	148	84	AAR10473	494	280	1.00E-74	175	147	84
AAR10738	506	155	6.00E-37	100	81	81	AAR10473	494	153	2.00E-31	100	80	80
AAR10731	500	280	9.00E-75	175	148	84	AAR10472	494	280	1.00E-74	175	147	84
AAR10731	500	152	4.00E-36	98	79	80	AAR10472	494	152	3.00E-36	98	79	80
AAR10725	496	280	9.00E-75	175	148	84	AAB33953	397	280	1.00E-74	276	169	61
AAR10725	496	155	4.00E-37	100	81	81	P52615	501	279	2.00E-74	175	147	84
AAR10723	496	280	9.00E-75	175	148	84	P52615	501	155	4.00E-37	100	81	81
AAR10723	496	153	2.00E-36	100	80	80	AAT81645	496	279	3.00E-74	175	147	84
AAR10722	496	280	9.00E-75	175	148	84	AAT81645	496	153	2.00E-36	100	80	80
AAR10722	496	152	3.00E-36	98	79	80	AAP13304	456	279	3.00E-74	245	160	65
AAR10693	495	280	9.00E-75	175	148	84	AAP13304	456	142	3.00E-33	90	73	81
AAR10693	495	154	7.00E-37	100	81	81	AAR10732	500	279	3.00E-74	175	147	84
AAR10692	495	280	9.00E-75	175	148	84	AAR10732	500	153	2.00E-36	100	80	80
AAR10692	495	152	4.00E-36	98	79	80	AAR10685	499	279	3.00E-74	175	147	84
AAR10684	499	280	9.00E-75	175	148	84	AAR10685	499	152	4.00E-36	98	79	80

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#ident	%S
AAR10477	506	279	3.00E-74	232	158	68	AAD28528	555	254	5.00E-67	248	143	57
AAR10477	506	155	4.00E-37	100	81	81	AAD28528	555	129	3.00E-29	108	67	63
AAR10470	497	279	3.00E-74	175	147	84	ZP_00091764573	254	9.00E-67	262	144	54	
AAR10470	497	155	3.00E-37	273	112	41	ZP_00091764573	139	4.00E-32	90	71	78	
YP_150203	495	278	3.00E-74	175	147	84	AAT06391	468	253	2.00E-66	158	134	84
YP_150203	495	156	2.00E-37	100	82	82	AAT06391	468	123	2.00E-27	117	70	59
AAR10582	501	278	3.00E-74	175	147	84	AAP13314	461	251	6.00E-66	177	133	75
AAR10582	501	152	4.00E-36	98	79	80	AAP13314	461	118	8.00E-26	280	93	33
AAR10556	495	278	3.00E-74	175	147	84	AAC45643	303	250	1.00E-65	303	140	46
AAR10556	495	155	6.00E-37	100	81	81	ZP_00173465285	249	2.00E-65	297	145	48	
IUCU	494	278	3.00E-74	174	147	84	AAT06390	460	249	2.00E-65	158	132	83
IUCU	494	150	1.00E-35	98	78	79	AAT06390	460	123	2.00E-27	189	82	43
AAB33952	405	278	3.00E-74	174	147	84	AAP13315	399	249	3.00E-65	158	127	80
S15121	494	278	3.00E-74	174	147	84	AAP13315	399	107	1.00E-22	304	93	30
S15121	494	152	4.00E-36	98	79	80	AAQ61541	282	248	6.00E-65	297	137	46
AAP13301	461	278	6.00E-74	244	157	64	AAL16054	455	248	6.00E-65	157	130	82
AAP13301	461	142	3.00E-33	90	73	81	AAL16054	455	116	3.00E-25	78	59	75
AAR10649	495	278	6.00E-74	175	147	84	ZP_00288126272	247	8.00E-65	296	145	48	
AAR10649	495	152	4.00E-36	98	79	80	AAQ61540	282	246	2.00E-64	297	137	46
A24262	490	277	7.00E-74	175	146	83	AAC72199	277	243	2.00E-63	292	136	46
A24262	490	153	2.00E-36	100	80	80	AAF19180	305	242	3.00E-63	304	141	46
AAD28524	556	276	2.00E-73	258	154	59	AAC45642	305	239	3.00E-62	304	139	45
AAD28524	556	123	2.00E-27	103	64	62	ZP_00289013269	238	5.00E-62	292	133	45	
AAD28521	560	276	2.00E-73	258	154	59	AAF19179	305	238	7.00E-62	305	142	46
AAD28521	560	132	5.00E-30	107	68	63	ZP_00149772273	236	1.00E-61	298	131	44	
AAN62540	416	275	3.00E-73	222	151	68	AAC79723	281	236	2.00E-61	295	129	43
AAN62540	416	143	2.00E-33	335	118	35	ZP_00289014271	236	2.00E-61	295	133	44	
AAP13317	455	275	3.00E-73	238	159	66	NP_718792	272	236	2.00E-61	297	140	47
AAP13317	455	140	1.00E-32	90	72	80	ZP_00274388386	236	2.00E-61	167	122	73	
AAN77106	449	275	5.00E-73	243	158	65	ZP_00274388386	135	6.00E-31	92	68	73	
AAN77106	449	126	3.00E-28	85	65	76	ZP_00289011271	235	3.00E-61	296	133	44	
AAP13312	460	274	6.00E-73	158	141	89	ZP_00288132271	235	4.00E-61	295	136	46	
AAP13312	460	142	3.00E-33	90	73	81	ZP_00288131271	234	6.00E-61	295	136	46	
AAN77105	454	274	6.00E-73	242	155	64	ZP_00299645314	234	7.00E-61	296	137	46	
AAN77105	454	126	3.00E-28	85	65	76	ZP_00288133271	233	1.00E-60	295	135	45	
AAP13308	502	273	1.00E-72	281	166	59	YP_149894	297	233	2.00E-60	303	133	43
AAP13308	502	144	1.00E-33	88	73	82	NP_718793	273	233	2.00E-60	297	138	46
AAD28523	565	273	2.00E-72	208	143	68	BAD06421	282	233	2.00E-60	299	137	45
AAD28523	565	117	2.00E-25	99	60	60	AAC72198	278	232	3.00E-60	296	137	46
NP_883763	392	272	2.00E-72	392	180	45	AAL57341	283	231	8.00E-60	300	138	46
AAN77109	455	272	2.00E-72	158	140	88	ZP_00288136271	230	1.00E-59	298	132	44	
AAN77109	455	130	1.00E-29	294	104	35	AAK14995	348	230	1.00E-59	348	157	45
AAR10726	499	272	2.00E-72	197	149	75	AAK14997	350	230	1.00E-59	350	154	44
AAR10726	499	152	4.00E-36	98	79	80	AAT86134	274	229	2.00E-59	289	127	43
AAD28519	523	272	3.00E-72	251	156	62	ZP_00211492272	229	2.00E-59	291	134	46	
AAD28519	523	128	7.00E-29	93	62	66	AAT86133	274	229	2.00E-59	289	127	43
AAN77108	448	271	5.00E-72	235	157	66	ZP_00289022272	228	4.00E-59	296	136	45	
AAN77108	448	121	9.00E-27	85	62	72	NP_954080	276	228	5.00E-59	298	133	44
AAN77107	452	270	1.00E-71	155	139	88	BAD06420	282	227	9.00E-59	299	131	43
AAN77107	452	124	1.00E-27	84	64	76	BAD01155	282	226	2.00E-58	299	131	43
AAD28529	555	268	5.00E-71	253	151	59	NP_791772	282	223	1.00E-57	297	129	43
AAD28529	555	132	5.00E-30	107	68	63	AAG15516	281	222	3.00E-57	291	129	44
NP_879790	391	268	6.00E-71	391	182	46	BAD06419	282	222	4.00E-57	297	129	43
NP_880978	391	266	2.00E-70	391	183	46	BAC54116	116	220	1.00E-56	116	116	100
AAD28525	560	265	4.00E-70	204	140	68	NP_522175	276	220	1.00E-56	296	134	45
AAD28525	560	114	1.00E-24	97	58	59	ZP_00302555327	219	2.00E-56	329	133	40	
AAD28518	550	265	4.00E-70	254	149	58	NP_871067	421	219	2.00E-56	181	109	60
AAD28518	550	119	3.00E-26	101	63	62	NP_871067	421	120	2.00E-26	345	109	31
AAD28526	508	265	5.00E-70	247	153	61	AAD46086	291	219	2.00E-56	292	124	42
AAD28526	508	112	5.00E-24	82	53	64	AAG15515	290	219	3.00E-56	292	123	42
AAO65441	466	263	1.00E-69	160	137	85	ZP_00127282271	218	4.00E-56	283	128	45	
AAO65441	466	138	7.00E-32	161	84	52	BAB58972	333	217	9.00E-56	330	134	40
AAD28527	517	262	3.00E-69	245	152	62	YP_080862	310	215	5.00E-55	313	135	43
AAD28527	517	126	2.00E-28	93	62	66	S69767	378	215	5.00E-55	307	147	47
AAD28520	521	259	3.00E-68	246	151	61	NP_244483	272	214	1.00E-54	294	124	42
AAD28520	521	121	9.00E-27	90	58	64	AAK14996	348	213	2.00E-54	273	137	50
AAL30512	460	254	5.00E-67	160	133	83	ZP_00263336255	213	2.00E-54	269	125	46	
AAL30512	460	92.4	4.00E-18	64	46	71	AAF09167	290	211	5.00E-54	300	125	41

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AAC38159	384	211	9.00E-64	381	141	37	ZP_00200889296	184	7.00E-46	300	110	36	
ZP_00183380275	210	1.00E-53	299	128	42		YP_129125	382	184	9.00E-46	382	130	34
NP_782313	275	2.00E-53	296	125	42		Q86572	376	184	9.00E-46	375	128	34
NP_693649	338	209	2.00E-53	338	131	36	BAB16757	492	184	9.00E-46	235	116	49
ZP_00289017266	209	2.00E-53	285	125	43	BAB16757	492	114	1.00E-24	261	80	31	
ZP_00289012373	209	3.00E-53	282	117	41	BAB58979	261	184	9.00E-46	270	111	41	
AAD12054	383	209	3.00E-53	380	141	37	AAC65757	288	183	2.00E-45	298	114	38
NP_391416	304	208	6.00E-53	306	130	42	AAC27808	377	183	2.00E-45	376	125	33
BAB87734	270	207	7.00E-53	294	123	41	AAQ98738	228	182	3.00E-45	251	102	40
NP_301058	284	207	9.00E-53	301	119	39	NP_348182	278	182	3.00E-45	292	115	39
AAD19738	279	207	9.00E-53	299	120	40	YP_010660	298	182	3.00E-45	311	119	38
NP_348820	275	207	1.00E-52	295	122	41	BAC98360	384	182	3.00E-45	381	132	34
YP_162364	284	206	2.00E-52	296	119	40	AAF95287	376	182	4.00E-45	375	133	35
AAB07350	284	206	3.00E-52	301	119	39	CAG25467	277	182	4.00E-45	292	107	36
NP_795640	377	202	2.00E-51	376	132	35	CAA45382	285	181	6.00E-45	301	112	37
ZP_00313278273	201	9.00E-51	295	120	40	ZP_00130180299	181	6.00E-45	285	107	36		
ZP_00236467307	200	1.00E-50	309	119	38	BAB16756	492	181	6.00E-45	236	111	47	
BAB87735	263	199	2.00E-50	287	118	41	BAB16756	492	115	6.00E-25	284	83	29
NP_967402	282	199	3.00E-50	295	124	42	AAC65834	266	181	9.00E-45	304	107	35
NP_967404	277	198	3.00E-49	292	120	41	AAQ98739	228	181	9.00E-45	251	105	41
NP_542301	389	196	3.00E-49	212	113	53	AAC65935	285	181	9.00E-45	301	111	36
NP_542301	389	103	1.00E-21	307	87	28	P21989	286	180	1.00E-44	304	113	37
ZP_00238470278	196	3.00E-49	293	117	39	AAQ98740	228	180	1.00E-44	251	105	41	
ZP_00236469286	195	4.00E-49	292	118	40	AAP08836	273	189	2.00E-44	298	115	38	
ZP_00244371402	195	5.00E-49	400	144	36	BAC98370	384	180	2.00E-44	381	131	34	
BAB58986	308	195	5.00E-49	302	118	39	CAG25466	260	180	2.00E-44	296	107	36
AAC31966	383	194	6.00E-49	380	136	35	AAA68044	290	179	2.00E-44	304	118	38
AAD24680	382	194	8.00E-49	379	136	35	JG6021	379	179	2.00E-44	379	129	34
AAD24678	382	194	8.00E-49	379	136	35	YP_123618	475	179	2.00E-44	309	124	40
CAG38109	277	194	8.00E-49	291	119	40	YP_123618	475	110	1.00E-23	209	77	38
ZP_0031275272	193	1.00E-48	292	118	40	BAC98368	384	179	2.00E-44	381	131	34	
ZP_00338286069	193	1.00E-48	290	120	41	S24986	288	179	3.00E-44	296	107	36	
ZP_00338286069	114	1.00E-24	169	67	42	YP_005879	367	179	3.00E-44	365	125	34	
NP_637306	399	193	1.00E-48	241	119	43	AAN86121	328	179	4.00E-44	335	120	36
NP_637306	399	104	1.00E-21	319	93	29	AAK20919	279	179	4.00E-44	296	109	36
ZP_00244372404	193	1.00E-48	404	150	37	AAK57644	281	179	4.00E-44	301	109	36	
AAS91569	379	193	1.00E-48	379	143	36	YP_095369	475	178	5.00E-44	309	125	40
BAB58978	306	193	1.00E-48	302	117	38	YP_095369	475	109	3.00E-23	198	78	39
BAB87737	266	192	2.00E-48	287	119	41	CAD42896	475	178	5.00E-44	309	123	39
YP_086397	1128	192	3.00E-48	252	120	47	CAD42866	475	109	3.00E-23	198	78	39
YP_065397	1128	107	2.00E-22	254	79	31	BAAB82629	336	178	5.00E-44	342	116	33
YP_001491	286	192	4.00E-48	302	112	37	YP_001839	281	177	1.00E-43	308	107	35
NP_712598	285	192	4.00E-48	302	112	37	NP_971613	286	177	1.00E-43	304	109	35
BAB58984	308	191	5.00E-48	304	119	39	BAAB8233	336	177	1.00E-43	342	115	33
BAB58980	308	191	7.00E-48	304	119	39	AAQ60166	265	177	1.00E-43	299	104	34
B32019	293	190	2.00E-47	304	121	39	YP_066429	857	177	1.00E-43	231	108	48
BAB58983	308	190	2.00E-47	304	119	39	YP_066429	857	115	5.00E-26	227	81	36
AAF95333	379	189	2.00E-47	375	130	34	YP_011656	297	177	1.00E-43	310	113	36
YP_109915	388	189	2.00E-47	385	136	35	CAA49317	335	177	1.00E-43	342	114	33
AAP08635	272	189	3.00E-47	296	114	38	YP_126643	475	176	2.00E-43	309	122	39
AAD27808	387	189	3.00E-47	334	136	35	YP_126643	475	110	2.00E-23	251	88	35
II01	398	189	3.00E-47	122	98	80	YP_001490	282	176	2.00E-43	300	107	35
II01	398	724	5.00E-12	54	36	66	CAD43143	475	176	2.00E-43	309	122	39
AAC71051	383	189	3.00E-47	335	134	34	CAD43143	475	110	2.00E-23	251	88	35
YP_076801	275	189	3.00E-47	298	114	38	CAD42899	327	176	2.00E-43	309	122	39
NP_967579	277	188	5.00E-47	301	117	38	CAD42894	361	176	2.00E-43	309	122	39
AAD24677	387	188	5.00E-47	334	136	35	CAA68234	475	178	2.00E-43	309	122	39
YP_027849	287	188	6.00E-47	232	112	38	CAA68234	475	109	3.00E-23	251	87	34
CAB67250	285	187	8.00E-47	300	111	37	CAD42900	489	176	2.00E-43	309	122	39
AAD24679	387	187	8.00E-47	334	135	35	CAD42900	489	110	2.00E-23	251	88	35
AAC01552	379	187	8.00E-47	375	129	34	BAA82930	336	176	2.00E-43	342	115	33
NP_972081	285	187	1.00E-46	300	113	37	NP_970090	277	176	2.00E-43	298	110	37
ZP_00236468284	187	1.00E-46	295	111	37	AAF95288	378	176	2.00E-43	377	134	35	
BAAB98371	384	187	1.00E-46	382	131	34	AAQ01655	378	175	2.00E-43	377	134	35
ZP_00288129272	186	2.00E-46	296	112	37	AAU07005	335	176	3.00E-43	342	113	33	
NP_967577	274	186	3.00E-46	292	115	39	NP_212281	336	176	3.00E-43	342	113	33
CAAA47610	276	186	5.00E-46	292	107	36	NP_712599	262	176	3.00E-43	300	107	35
BAAB98982	249	185	5.00E-46	269	114	42	CAA49307	336	176	3.00E-43	342	114	33

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
BAB50981	280	176	3.00E-43	285	113	39	AAA99807	488	168	5.00E-41	242	104	42
CAA53013	336	175	4.00E-43	342	114	33	AAA99807	488	113	2.00E-24	335	100	29
BAA07832	334	175	4.00E-43	337	115	34	AAR00324	336	168	8.00E-41	342	110	32
CAA33695	335	175	4.00E-43	342	113	33	BAA11604	329	158	6.00E-41	335	108	32
CAA37884	334	175	4.00E-43	340	116	34	AAQ6080	372	167	8.00E-41	369	122	33
NP_521943	273	175	4.00E-43	297	109	36	BAA11603	329	167	8.00E-41	335	108	32
AAF95332	377	175	4.00E-43	377	129	34	BAA11597	329	167	8.00E-41	335	109	32
CAA59172	474	175	4.00E-43	228	105	46	BAA82631	329	167	8.00E-41	335	109	32
CAA59172	474	107	1.00E-22	214	79	38	BAA11611	327	167	1.00E-40	333	109	32
AAB37005	336	175	4.00E-43	342	113	33	NP_249783	488	167	1.00E-40	242	103	42
CAA49316	336	175	5.00E-43	342	113	33	NP_249783	488	113	2.00E-24	335	100	29
AAF87586	273	175	5.00E-43	300	111	37	P21184	394	167	1.00E-40	271	105	38
AAG34566	280	175	5.00E-43	294	107	36	ZP_00138680488	167	1.00E-40	242	103	42	
CAA53012	334	174	7.00E-43	333	113	33	ZP_00138680488	113	2.00E-24	335	108	29	
CAD42893	493	174	7.00E-43	310	122	39	ZP_00130923288	167	1.00E-40	284	103	36	
CAD42893	493	69.8	3.00E-20	191	72	37	AAC63946	390	167	1.00E-40	271	105	38
CAA49308	336	174	7.00E-43	342	113	33	AAP33170	384	167	1.00E-40	271	105	38
AAS91572	375	174	7.00E-43	370	126	34	BAA11607	329	167	1.00E-40	335	108	32
AAS91571	400	174	7.00E-43	377	126	33	AAC27900	378	167	1.00E-40	377	126	33
CAA53014	336	174	9.00E-43	342	113	33	NP_798837	376	167	1.00E-40	377	125	33
AAK01040	282	174	9.00E-43	300	107	36	NP_798838	378	167	1.00E-40	377	126	33
CAB67249	286	174	1.00E-42	304	103	33	AAC63950	478	167	1.00E-40	225	101	44
CAA49322	336	174	1.00E-42	342	112	32	AAC63950	478	111	7.00E-24	278	83	29
CAA49318	336	174	1.00E-42	342	113	33	BAA09687	329	167	1.00E-40	335	107	31
BAA82635	323	174	1.00E-42	329	114	34	BAA09681	329	167	1.00E-40	336	108	32
CAA34735	336	173	2.00E-42	342	112	32	BAA11500	329	167	1.00E-40	335	107	31
YP_013380	287	173	2.00E-42	296	107	36	BAA82636	324	167	1.00E-40	323	109	33
CAA45011	336	173	2.00E-42	342	113	33	BAA09689	329	166	2.00E-40	335	107	31
NP_470041	287	173	2.00E-42	296	107	36	BAA09688	329	166	2.00E-40	335	107	31
NP_464217	287	173	2.00E-42	296	107	36	BAA09686	329	166	2.00E-40	335	107	31
NP_797167	384	173	2.00E-42	384	125	32	BAA09684	329	166	2.00E-40	335	107	31
AAQ10326	376	173	2.00E-42	375	121	32	BAA09678	329	166	2.00E-40	335	107	31
AAU95795	287	173	2.00E-42	296	107	36	BAA11610	327	166	2.00E-40	333	110	33
CAA49321	336	173	2.00E-42	342	113	33	A37853	394	166	3.00E-40	271	105	38
AAP33175	387	173	2.00E-42	314	118	37	A37853	394	107	2.00E-22	197	73	37
AAP33175	387	122	5.00E-27	182	73	40	AAD10272	378	165	5.00E-40	377	125	33
BAA82634	319	173	2.00E-42	323	113	34	AAG29830	327	165	5.00E-40	333	108	32
AAK57645	286	172	3.00E-42	301	109	36	AAP08537	249	165	5.00E-40	271	98	36
NP_969824	277	172	3.00E-42	295	109	36	AAB03518	335	165	5.00E-40	334	111	33
YP_061686	277	172	3.00E-42	299	106	35	ZP_00317212580	164	7.00E-40	222	101	45	
CAD42895	491	172	3.00E-42	166	92	55	ZP_00317212580	99.4	4.00E-20	182	66	36	
CAD42895	491	109	3.00E-23	198	78	39	AAA62843	688	164	9.00E-40	252	103	40
NP_972083	285	172	4.00E-42	304	106	34	AAA62843	688	95.9	4.00E-19	92	46	50
YP_158591	306	171	6.00E-42	151	89	58	NP_746492	687	164	9.00E-40	252	103	40
BAB65985	280	171	6.00E-42	280	110	39	NP_746492	687	96.3	3.00E-19	129	53	41
AAR16426	329	171	7.00E-42	314	117	37	BAA33017	318	164	1.00E-39	323	108	33
AAQ10324	377	171	7.00E-42	376	123	32	ZP_00100062385	163	2.00E-39	385	128	33	
ZP_00346385295	171	7.00E-42	292	98	33	AAC99335	283	163	2.00E-39	301	100	33	
AAC28557	387	171	7.00E-42	314	117	37	ZP_00317210587	162	5.00E-39	254	105	41	
BAA11613	326	171	7.00E-42	326	109	33	ZP_00317210587	95.9	4.00E-19	129	55	42	
BAA26807	320	171	7.00E-42	323	111	34	NP_712200	283	162	5.00E-39	301	100	33
NP_348785	283	171	1.00E-41	290	104	35	CAI665866	268	162	5.00E-39	103	84	81
AAU65734	287	171	1.00E-41	236	106	35	YP_001838	283	161	6.00E-39	301	99	32
AAP33174	387	171	1.00E-41	314	117	37	AAC99335	283	161	8.00E-39	301	99	32
AAP33174	387	122	5.00E-27	182	73	40	AAC63947	393	160	1.00E-38	300	108	36
AAC01556	377	171	1.00E-41	379	127	33	NP_770335	274	160	1.00E-38	295	95	32
BAA28506	323	171	1.00E-41	323	111	34	AAB94024	283	160	2.00E-38	301	98	32
NP_782278	280	170	1.00E-41	294	105	35	AAC63948	486	160	2.00E-38	232	99	42
ZP_00279040381	170	1.00E-41	378	127	33	AAC63948	486	100	1.00E-20	346	89	25	
CAA53011	336	170	1.00E-41	342	111	32	CAA45081	285	159	2.00E-38	304	108	35
BAA11602	329	170	1.00E-41	335	111	33	BAA23224	328	159	3.00E-38	327	106	32
AAF95268	377	169	2.00E-41	379	127	33	CAC36206	274	159	3.00E-38	273	94	34
YP_011295	297	169	2.00E-41	295	101	34	BAB13814	413	159	3.00E-38	256	101	39
BAA11505	329	169	2.00E-41	335	110	32	BAB13814	413	89.7	3.00E-17	173	60	34
CAA53015	336	169	3.00E-41	334	112	33	BAA11602	328	159	3.00E-38	327	108	32
CAD42887	476	169	3.00E-41	314	122	38	AAC63949	669	159	4.00E-38	260	105	40
CAD42897	476	109	3.00E-23	198	78	39	AAC63949	669	106	5.00E-23	273	74	27
AAC28566	387	169	4.00E-41	300	114	38	AAR08139	236	158	5.00E-38	217	92	42

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
CAC36204	274	158	5.00E-38	273	95	34	YP_035878	268	142	5.00E-33	292	97	33
AAC99334	283	158	5.00E-38	301	98	32	BAA12719	292	142	5.00E-33	291	92	31
AAU93692	286	158	7.00E-38	304	106	34	BAA20904	289	142	5.00E-33	291	92	31
BAA82632	311	158	7.00E-38	313	103	32	BAA19430	292	142	5.00E-33	291	92	31
ZP_00218989614	158	2.00E-37	242	97	40	AAO08752	374	141	6.00E-33	167	76	45	
ZP_00218989614	90.1	2.00E-17	86	46	53	NP_782259	268	141	6.00E-33	266	79	29	
BAB87728	413	156	2.00E-37	278	102	36	BAD12778	248	141	6E-33	267	88	32
BAB87728	413	85.1	7.00E-16	108	45	141	292	141	6.00E-33	291	92	31	
CAC36205	274	156	2.00E-37	273	93	34	BAA20481	292	141	6.00E-33	291	92	31
AAC38200	504	156	2.00E-37	246	97	39	BAA20479	292	141	6.00E-33	291	92	31
AAC38200	504	92	6.00E-18	87	45	52	BAA20477	292	141	6.00E-33	291	92	31
BAB87730	375	155	3.00E-37	374	115	30	BAA13582	292	141	6.00E-33	291	92	31
AAB09433	377	155	6.00E-37	182	80	43	BAA19444	292	141	6.00E-33	291	92	31
ZP_00270146298	154	1.00E-36	305	107	35	BAA19443	292	141	6.00E-33	291	92	31	
NP_935285	396	153	2.00E-36	376	118	31	BAA19439	292	141	6.00E-33	291	92	31
AA91570	377	153	2.00E-36	218	89	40	BAA89303	248	141	6.00E-33	267	87	32
CAC36223	271	152	3.00E-36	268	91	33	BAA89302	248	141	6.00E-33	267	88	32
NP_945993	274	152	4.00E-36	295	92	31	BAA77944	248	141	6.00E-33	267	87	32
AA008751	377	151	6.00E-36	376	120	31	BAC16349	248	141	8.00E-33	267	87	32
BAB87729	384	151	6.00E-36	169	82	48	BAA77321	208	141	8.00E-33	214	87	40
AA061408	241	151	8.00E-36	253	98	36	AAQ81414	241	140	1.00E-32	267	92	34
AA010325	377	151	8.00E-36	376	119	31	ZP_00004597493	140	1.00E-32	166	74	44	
YP_128298	426	151	8.00E-36	251	106	42	ZP_00004597493	97.1	2.00E-19	176	63	35	
YP_128298	426	89.4	4.00E-17	149	55	36	YP_D83129	266	140	1.00E-32	292	95	32
BAB87732	377	151	8.00E-36	169	82	48	CAA75001	493	140	1.00E-32	166	74	44
BAB87732	377	79.7	3.00E-14	291	83	28	CAA75001	493	97.1	2.00E-19	176	63	35
AAB09434	377	150	1.00E-35	376	119	31	AAG21990	170	140	1.00E-32	166	74	44
AAB03521	334	150	1.00E-35	335	102	30	BAA20478	292	140	1.00E-32	291	92	31
AAC38201	540	150	2.00E-35	266	100	37	AAQ8634	266	140	1.00E-32	291	95	32
AAC38201	540	51.2	1.00E-05	147	43	29	BAD13295	248	140	1.00E-32	267	87	32
ZP_00329901380	149	2.00E-35	382	125	32	AAO61411	241	140	2.00E-32	267	92	34	
NP_935286	377	149	2.00E-35	376	112	29	NP_228567	387	140	2.00E-32	392	119	30
Q56574	377	149	2.00E-35	376	115	30	AAK31137	257	140	2.00E-32	273	87	31
AA061409	241	149	3.00E-35	253	95	36	BAAB9304	248	140	2.00E-32	267	87	32
NP_933764	385	149	4.00E-35	178	80	44	BAA77325	200	139	3.00E-32	218	89	40
NP_933764	365	102	6.00E-21	301	92	30	AAQ96735	620	139	4.00E-32	258	96	37
CAC36218	220	149	4.00E-35	238	90	37	AAQ96735	620	63.2	3.00E-09	64	31	48
BAA15744	340	149	4.00E-35	151	80	52	AAO61415	240	139	4.00E-32	266	91	34
BAA77322	201	149	4.00E-35	214	90	42	CAC36207	234	139	4.00E-32	248	85	34
YP_165525	471	148	5.00E-35	170	82	48	ZP_00236466266	138	5.00E-32	292	96	32	
YP_165525	471	99.4	4.00E-20	267	77	28	BAB87738	554	138	7.00E-32	136	71	52
AAD10271	384	147	9.00E-35	188	78	41	BAB87738	554	99.8	3.00E-20	291	82	28
CAC36221	222	147	1.00E-34	237	85	35	YP_083130	450	137	9.00E-32	289	96	33
AA061406	241	147	2.00E-34	267	96	35	YP_083130	450	80.9	1.00E-14	111	46	41
CAC36220	222	147	2.00E-34	237	84	35	NP_348261	239	137	1.00E-31	292	89	30
ZP_00273972629	146	3.00E-34	199	85	42	AAO61412	240	135	3.00E-31	266	90	33	
ZP_00273972629	104	9.00E-22	92	57	61	AAQ96736	470	134	8.00E-31	233	86	36	
YP_129126	393	145	4.00E-34	213	82	38	AAQ96736	470	65.1	8.00E-10	264	68	25
YP_129126	393	90.5	2.00E-17	203	63	31	BAA89305	248	134	1.00E-30	268	84	31
NP_978099	266	145	4.00E-34	292	96	32	CAC36202	234	132	5E-30	248	83	33
AAO61410	241	145	6.00E-34	263	91	34	AAT06255	459	132	5.00E-30	139	70	50
AAB03520	334	145	6.00E-34	330	102	30	AAT06255	459	87.4	1E-16	159	57	35
BAA77320	202	145	6.00E-34	217	91	41	YP_050963	484	131	7.00E-30	129	71	56
AA061407	241	144	8.00E-34	264	93	35	YP_050963	484	88.6	6.00E-17	152	63	41
CA44444	377	144	8.00E-34	377	112	29	AAB82610	472	131	7.00E-30	265	93	35
CAAA67103	333	144	1.00E-33	333	108	32	AAB82610	472	98.2	8.00E-20	153	63	41
CAC36219	220	144	1.00E-33	238	89	37	CAC36217	223	131	9.00E-30	241	78	32
AAQ96734	625	143	2.00E-33	263	99	37	AAA62844	479	130	1.00E-29	132	69	52
AAQ96734	626	70.9	1.00E-11	68	36	AAA62844	479	97.4	1.00E-19	260	77	27	
BAA12718	292	143	2.00E-33	291	94	32	BAA77323	200	130	1.00E-29	216	85	39
AAO61404	241	143	2.00E-33	264	92	34	NP_242343	464	130	2.00E-29	135	70	51
AAO61405	241	142	3.00E-33	264	89	33	NP_242343	464	95.9	4.00E-19	169	57	33
BAA12102	292	142	3.00E-33	291	94	32	CAC36201	223	129	4.00E-29	241	79	32
BAA13584	292	142	3.00E-33	291	93	31	AAT06254	493	129	4.00E-29	140	68	48
YP_148995	604	142	4.00E-33	269	96	35	AAT06254	493	94	2.00E-18	195	67	34
YP_148995	604	103	3.00E-21	179	62	34	AAB82613	508	128	7.00E-29	150	68	45
CAC36224	222	142	4.00E-33	237	86	36	AAB82613	508	86.7	2.00E-16	158	58	36
ZP_00267990272	142	5.00E-33	280	85	30	NP_978100	465	126	2.00E-28	221	77	34	

Fig. 25B-Continued

Hit_ID	Length	Blitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Blitscore	Expected	Length	#Ident	%S
NP_978100	465	76.1	7.00E-13	84	41	48	BAA77299	398	55.8	5.00E-07	130	43	33
AAL26808	229	126	2.00E-28	248	79	31	CAA67104	343	112	3.00E-24	225	78	33
AAQ96737	430	124	1.00E-27	165	67	40	CAA67104	349	107	1.00E-22	187	70	37
AAQ96737	430	60.1	2.00E-08	92	33	35	CAA67101	434	112	5.00E-24	195	75	38
CAB46359	81	124	1E-27	81	63	77	CAA67101	434	98.2	8.00E-20	327	91	27
CAC36222	223	122	6.00E-27	240	77	32	BAA77312	170	111	7.00E-24	125	61	48
AAP34182	184	122	5.00E-27	182	73	40	BAA77310	153	111	7.00E-24	125	61	48
AAP34178	185	122	5.00E-27	182	73	40	CAA66928	434	110	1.00E-23	196	74	37
AAO08750	375	121	7.00E-27	374	108	28	CAA66928	434	91.7	8.00E-18	322	86	26
AAP44578	259	121	7.00E-27	260	83	31	BAA77302	176	110	1.00E-23	174	74	42
NP_933768	375	121	7.00E-27	374	108	28	BAD22847	568	110	2.00E-23	180	64	35
AAC09389	259	119	3.00E-26	195	77	38	BAD22847	568	75.6	6.00E-13	83	34	40
AAP34194	195	119	3.00E-26	195	77	39	NP_797170	374	109	3.00E-23	148	60	40
AAP34185	194	119	3.00E-26	195	77	39	NP_797170	374	74.3	1.00E-12	217	61	28
CAA67102	434	119	3.00E-26	197	78	39	NP_903288	513	109	3.00E-23	294	89	30
CAA67102	434	98.1	7.00E-19	310	91	29	NP_903288	513	73.2	3.00E-12	122	44	36
AAQ03611	128	118	6.00E-26	117	60	51	BAD22846	568	109	3.00E-23	180	64	35
CAA66927	434	118	6.00E-26	195	78	39	BAD22846	568	75.5	8.00E-13	83	34	40
CAA66927	434	101	1.00E-20	327	92	28	BAD22839	568	109	3.00E-23	180	63	35
AAG03603	127	118	8.00E-26	120	59	49	BAD22839	568	74.7	1.00E-12	83	34	40
BAD22842	567	118	8.00E-26	180	68	37	AAK12380	576	109	3.00E-23	184	63	34
BAD22842	567	75.9	4.00E-13	84	34	40	AAK12380	576	79	5.00E-14	107	43	40
BAA77307	483	118	8.00E-26	254	87	34	AAK12377	574	109	3.00E-23	184	63	34
BAA77307	483	68.5	7.00E-08	221	60	27	AAK12377	574	79.3	4.00E-14	124	47	37
AAQ03617	132	117	1.00E-25	117	59	50	AAF25214	576	109	3.00E-23	184	63	34
AAQ03618	133	117	1.00E-25	117	59	50	AAF25214	576	77	2.00E-13	83	37	44
AAQ03614	131	117	1.00E-25	117	59	50	AAC25644	578	109	3.00E-23	184	62	33
AAQ03610	129	117	1.00E-25	117	59	50	AAC25644	576	77.4	1.00E-13	156	52	33
BAD22849	566	117	2.00E-25	180	68	37	S41310	575	109	3.00E-23	182	68	37
BAD22849	566	75.5	6.00E-13	83	34	40	S41310	575	79.3	4.00E-14	107	43	40
BAD22843	567	117	2.00E-25	180	68	37	CAC03725	487	108	6.00E-23	144	65	45
BAD22843	567	74.7	1.00E-12	83	34	40	CAC03725	457	101	1.00E-20	276	85	30
BAD22841	557	117	2.00E-25	180	68	37	A42474	573	108	6.00E-23	184	63	34
BAD22841	567	74.7	1.00E-12	83	34	40	A42474	573	79.7	3.00E-14	124	46	37
BAD22837	567	117	2.00E-25	180	68	37	A39228	576	108	6.00E-23	184	62	33
BAD22837	567	74.7	1.00E-12	83	34	40	A39228	576	77	2.00E-13	83	37	44
BAD22835	566	117	2.00E-25	180	68	37	BAD22844	568	108	6.00E-23	180	63	35
BAD22835	566	75.5	6.00E-13	83	34	40	BAD22844	568	75.5	6.00E-13	83	34	40
BAD22834	566	117	2.00E-25	180	68	37	BAC44988	571	108	6.00E-23	181	66	36
BAD22834	566	75.5	6.00E-13	83	34	40	BAC44988	571	72	6.00E-12	83	34	40
CAC36203	221	116	2.00E-25	238	74	31	BAC44988	575	108	6.00E-23	184	63	34
BAD22836	567	116	2.00E-25	180	68	37	BAC44988	575	77.4	1.00E-13	159	55	34
BAD22836	567	74.7	1.00E-12	83	34	40	AAF05902	574	108	6.00E-23	184	63	34
NP_381395	160	116	3.00E-25	179	68	37	AAF05902	574	77	2.00E-13	83	37	44
BAD22838	563	116	3.00E-25	180	67	37	AAN75633	196	108	6.00E-23	215	72	33
BAD22838	563	76.6	3.00E-13	173	50	28	AAC25648	576	108	6.00E-23	184	62	33
BAD22845	565	115	4.00E-25	180	67	37	AAC25648	576	77.4	1.00E-13	83	37	44
BAD22845	565	78.2	9.00E-14	127	45	35	AAC25643	576	108	6.00E-23	184	63	34
AAB69135	434	115	4E-25	186	76	38	AAC25643	576	77.4	1.00E-13	124	44	35
AAB69135	434	103	2.00E-21	327	93	28	AAC25657	576	108	6.00E-23	184	62	33
AAB69134	434	115	4.00E-25	186	76	38	AAC25657	575	77	2.00E-13	83	37	44
AAB69134	434	102	4.00E-21	327	93	28	P27053	573	108	6.00E-23	184	63	34
I40573	195	115	4.00E-25	165	65	38	P27053	573	79.7	3.00E-14	124	46	37
BAD22840	566	115	5.00E-25	180	67	37	BAB63937	486	108	6.00E-23	183	67	36
BAD22840	566	76.5	6.00E-13	83	34	40	BAB63937	485	80.1	2.00E-14	83	38	45
AAQ03605	125	114	6E-25	127	61	48	AAF05902	574	77	2.00E-13	83	37	44
AAG61259	253	114	1.00E-24	257	80	31	NP_206915	514	108	8.00E-23	187	67	35
CAA67185	700	114	1.00E-24	301	93	30	NP_206915	514	77.4	1.00E-13	121	47	38
CAA67105	700	102	4.00E-21	298	92	30	BAB71798	486	108	8.00E-23	183	66	36
CAC03724	440	113	2.00E-24	237	83	35	BAB71798	486	86.1	2.00E-14	83	38	45
CAC03724	440	106	3.00E-22	255	80	31	AAP72265	514	107	1.00E-22	166	62	37
BAD22848	571	113	2E-24	180	66	36	AAP72265	514	77.4	1.00E-13	121	47	38
BAD22848	571	74.7	1.00E-12	83	33	84	BAC44987	574	107	1.00E-22	181	65	35
AAB66595	340	113	2E-24	225	76	33	BAC44987	574	82	8.00E-15	162	56	34
AAB66595	340	107	1E-22	187	70	37	BAC44986	574	107	1.00E-22	181	65	35
AAB66597	340	113	2.00E-24	225	76	33	BAC44986	574	79	5.00E-14	83	37	44
AAB66597	340	105	5.00E-22	187	89	36	AAD10273	374	107	1.00E-22	160	61	38
AAB66597	340	105	4.00E-22	215	86	40	AAD10273	374	76.9	4.00E-13	217	59	27
BAA77299	398	113	2.00E-24	215	86	40	AAP34188	183	107	1.00E-22	187	70	37

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AAC25639	575	107	1.00E-22	181	65	35	NP_420273	273	104	1.00E-21	297	76	25
AAC25639	575	82	6.00E-15	162	55	34	AAC25638	576	104	1.00E-21	181	64	35
AAA25016	514	107	1.00E-22	187	67	35	AAC25638	576	79.3	4.00E-14	121	44	36
AAA25016	514	77.4	1.00E-13	121	47	38	BAB85480	491	104	1.00E-21	181	63	34
NP_222828	514	107	1.00E-22	187	57	35	BAB85480	491	83.2	3.00E-15	174	55	31
NP_222828	514	77.4	1.00E-13	121	47	38	CAA82524	575	103	1.00E-21	182	69	37
AAL8694	514	107	1.00E-22	187	67	35	CAA82524	575	73.9	2.00E-12	107	40	37
AAL8694	514	77.4	1.00E-13	121	47	38	BAC56988	491	103	1.00E-21	184	63	34
CAA82523	575	107	1.00E-22	182	67	36	BAC56988	491	82.8	4.00E-15	174	59	33
CAA82523	575	79.3	4.00E-14	107	43	40	AAC25547	575	103	1.00E-21	171	62	36
CAB46859	514	107	1.00E-22	249	79	31	AAC25547	575	78.6	7.00E-14	107	43	40
CAB46859	514	75.1	7.00E-13	148	50	33	NP_907654	518	103	2.00E-21	192	63	32
AAN74969	514	107	1.00E-22	187	67	35	NP_907654	518	94.7	9.00E-19	333	85	25
AAN74969	514	77.4	1.00E-13	121	47	38	CAA72431	630	103	2.00E-21	181	64	35
BAC56997	575	107	1.00E-22	171	64	37	CAA72431	630	71.2	1.00E-11	105	38	36
BAC56997	575	79.7	3.00E-14	121	44	36	AAC25546	572	103	2.00E-21	181	67	37
AAA17046	575	107	1.00E-22	182	67	36	AAC25546	572	72.8	4.00E-12	132	45	34
AAA17046	575	73.9	2.00E-12	107	40	37	BAB71800	487	103	2.00E-21	183	63	34
CAA40460							BAB71800	487	80.1	2.00E-14	83	38	45
							BAC56991	491	103	3.00E-21	219	68	31
							BAC56991	491	84	2.00E-15	171	56	32
CAA40460	576	82	6.00E-15	162	56	34	AAC36988	273	103	3.00E-21	297	75	25
AAC25650	572	107	2.00E-22	171	64	37	NP_404368	404	102	3.00E-21	223	67	30
AAC25650	572	77.8	1E-13	132	48	36	NP_670739	401	102	3.00E-21	223	67	30
AAC25649	572	107	2.00E-22	181	65	35	NP_670739	401	57.4	2.00E-07	84	28	33
AAC25649	572	78.2	9.00E-14	132	48	36	AAV35033	543	102	4.00E-21	169	61	36
AAC25645	572	107	2.00E-22	181	65	35	AAV35033	543	62	6.00E-09	141	42	29
AAC25645	572	79.3	4.00E-14	132	48	36	BAC56990	490	102	4.00E-21	219	68	31
AAC25641	572	107	2.00E-22	181	65	35	BAC56990	490	83.6	2.00E-15	171	59	34
AAC25641	572	78.6	7.00E-14	83	37	44	BAAT7308	120	102	4.00E-21	119	55	46
BAC56996	572	106	2.00E-22	184	61	33	NP_282484	572	102	6.00E-21	181	66	36
BAC56996	572	79.7	3.00E-14	124	46	37	NP_282484	572	72.8	4.00E-12	132	45	34
NP_282485	572	106	3.00E-22	171	63	36	CAA72430	626	102	6.00E-21	183	65	35
NP_282485	572	79.3	4.00E-14	132	48	36	CAA72430	626	79.3	4.00E-14	148	50	33
BAA77318	125	106	3.00E-22	125	57	45	AAK73706	557	101	7.00E-21	178	59	33
BAA77318	131	106	3.00E-22	125	57	45	AAK73706	557	70.1	2.00E-11	113	41	36
BAC56994	487	105	4.00E-22	183	64	34	AAC25640	575	101	7.00E-21	181	65	35
BAC56994	487	80.1	2.00E-14	83	38	45	AAC25640	575	77.8	1.00E-13	296	78	26
BAA77304	165	105	4.00E-22	165	68	41	CAB46858	513	101	1.00E-20	240	73	30
B39228	576	105	5.00E-22	184	63	34	CAB46858	513	77.8	1.00E-13	329	89	27
B39228	576	71.6	8.00E-12	83	33	39	BAC56987	492	101	1.00E-20	174	61	35
AAB03519	328	105	5.00E-22	308	87	28	BAC56987	492	82.8	4.00E-15	175	58	33
AAF25216	576	105	5.00E-22	184	63	34	AAK73706	557	101	1.00E-20	178	59	33
AAF25216	576	73.6	2.00E-12	83	34	40	AAK73700	557	70.1	2.00E-11	113	41	36
B42474	573	105	5E-22	184	64	34	AAK73700	557	101	2.00E-11	113	41	36
B42474	573	73.9	2.00E-12	124	43	34	AAC25642	572	101	1.00E-20	181	66	36
A35146	573	105	5.00E-22	184	64	34	AAC25642	572	71.6	8.00E-12	132	45	34
A35146	573	72.8	4.00E-12	124	43	34	NP_214372	518	100	1.00E-20	209	62	29
P18245	573	105	5.00E-22	184	64	34	NP_214372	518	52	6.00E-09	129	39	30
P18245	573	73.9	2.00E-12	124	43	34	BAC56986	500	100	1.00E-20	153	61	39
BAA77314	124	105	5.00E-22	124	57	45	BAC56986	500	85.9	4.00E-16	316	82	25
AAA23020	573	105	5.00E-22	184	64	34	AAK73703	556	100	2.00E-20	175	62	35
AAA23020	573	72.8	4.00E-12	124	43	34	AAK73703	556	71.6	8.00E-12	287	73	25
1814303A	566	105	5.00E-22	182	66	36	AAK73702	558	100	2.00E-20	175	62	35
1814303A	566	45.4	6.00E-04	75	24	32	AAK73702	558	74.7	1.00E-12	287	72	25
AAM76286	576	105	7.00E-22	184	62	33	AAK73699	194	100	2.00E-20	175	62	35
AAM76286	576	73.6	2.00E-12	83	34	40	AAK73699	194	99.8	3.00E-20	115	58	50
B40566	514	104	9.00E-22	189	63	33	AAK73705	554	100	2.00E-20	175	61	34
B40566	514	80.5	2.00E-14	123	48	39	AAK73698	194	100	2.00E-20	175	61	34
AAP78393	514	104	9.00E-22	188	67	35	AAK73697	194	100	2.00E-20	175	61	34
AAP78393	514	68.9	5.00E-11	115	40	34	YP_061683	290	99.8	3.00E-20	308	85	27
BAC56995	487	104	9.00E-22	183	65	35	BAA77300	115	99.8	3.00E-20	115	58	50
BAC56995	487	80.1	2.00E-14	83	38	45	BAB71799	492	99.8	3.00E-20	174	59	33
BAC56992	487	104	9.00E-22	183	65	35	BAB71799	492	85.9	4.00E-16	172	57	33
BAC56992	487	79	5.00E-14	83	37	44	ZP_00313281302	99.4	4.00E-20	274	69	25	
BAC56989	487	104	9.00E-22	183	65	35	ZP_00100043955	99.4	4.00E-20	270	82	30	
BAC56989	487	79	5.00E-14	83	37	44	ZP_00100043955	63.5	2.00E-09	154	47	30	
BAB71801	491	104	9.00E-22	180	63	35	A44757	572	99	5.00E-20	171	58	33
BAB71801	491	81.6	8.00E-15	174	54	31	A44757	572	79.7	3.00E-14	124	46	37

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#ident	%S
NP_420274	273	99	5.00E-20	292	75	25	ZP_00339045281	84	2E-15	296	75	25	
NP_419609	273	99	5.00E-20	292	72	24	AAA85133	102	84	2E-15	87	43	49
AAA23321	572	99	5.00E-20	171	58	33	YP_080686	303	89.2	3E-15	281	70	24
AAA23021	572	79.7	3.00E-14	124	46	37	FLQL2C	276	82.4	5E-15	298	70	23
NP_419611	273	98.6	6.00E-20	292	72	24	NP_773505	314	80.9	1E-14	317	73	23
YP_071817	400	98.2	8.00E-20	207	61	29	YP_148989	297	80.5	2E-14	288	69	23
YP_071817	400	60.1	2.00E-08	115	37	32	NP_353570	313	79.7	3.00E-14	313	75	23
NP_865698	718	98.2	8.00E-20	284	88	30	ZP_00339044282	79.3	4.00E-14	298	72	24	
NP_865696	716	81.3	1.00E-14	143	57	39	CAD11203	461	78	5.00E-14	149	49	52
NP_622171	296	98.2	8.00E-20	290	83	28	CAD11203	461	63.9	2.00E-09	109	39	35
BAA77306	397	98.2	8.00E-20	188	57	35	CAD11202	461	78	5.00E-14	149	49	32
BAA77306	397	51.2	1.00E-05	91	27	29	CAD11202	461	63.9	2.00E-09	102	39	35
AAP78250	508	97.8	1.00E-19	170	60	35	ZP_00007767281	78.6	7.00E-14	302	70	23	
AAP78250	508	76.3	3.00E-13	297	74	24	NP_840399	299	77.8	1.00E-13	280	71	24
NP_419610	273	97.4	1.00E-19	292	70	23	CAD11205	458	77.8	1.00E-13	136	45	33
NP_868107	739	97.4	1.00E-19	257	76	29	CAD11208	458	67.8	1.00E-10	294	70	23
NP_868107	739	79.3	4.00E-14	250	70	28	CAD11207	458	77.8	1.00E-13	136	45	33
CAA43148	510	97.1	2.00E-19	170	57	33	CAD11207	458	65.9	4.00E-10	294	69	23
CAA43148	510	82.4	5.00E-15	311	78	25	CAD11208	458	77.8	1.00E-13	136	45	33
BAC86993	500	87.1	2.00E-19	232	73	32	CAD11205	458	65.9	4.00E-10	284	69	23
BAC86993	500	82.4	5.00E-15	174	65	31	ZP_00339046273	77.4	1.00E-13	280	73	26	
NP_223265	510	96.3	3.00E-19	170	57	33	ZP_00338034282	77.4	1.00E-13	294	72	24	
NP_223265	510	82.4	5.00E-15	311	78	25	AAAN0850139	139	77.4	1.00E-13	74	38	51
AAP72264	510	96.3	3.00E-19	170	57	33	CAC03722	433	77.4	1.00E-13	315	74	23
AAP72264	510	88.2	3.00E-15	311	78	25	CAC03722	433	31.8	9.30E+00	199	49	24
AAU21201	510	96.3	3.00E-19	170	57	33	BAB18984	209	77.4	1.00E-13	134	46	34
AAU21201	510	82	6.00E-15	311	78	25	AAM90027	145	77	2.00E-13	158	53	33
CAC03721	427	96.3	3.00E-19	251	76	30	BAC16347	200	77	2.00E-13	134	45	34
CAC03721	427	94.4	1.00E-18	271	74	27	BAC16341	200	77	2.00E-13	134	46	34
NP_404369	399	95.9	4.00E-19	295	76	25	BAC16340	200	77	2.00E-13	134	46	34
AAQ82717	510	95.9	4.00E-19	170	57	33	ZP_00193517283	76.6	3.00E-13	290	65	22	
AAQ82717	510	82	6.00E-15	311	78	25	AAR98327	69	76.6	3E-13	89	39	56
ZP_00299642294	95.5	5.00E-19	295	75	25	ZP_00337030282	75.3	3.00E-13	299	69	23		
A55864	518	95.5	5.00E-19	197	65	32	BAB19647	200	75.9	4.00E-13	134	45	33
A55864	518	94.7	9.00E-19	333	85	25	BAB18970	200	75.9	4.00E-13	134	45	33
AAB95381	273	95.5	5.00E-19	292	70	23	BAB18969	200	75.9	4.00E-13	134	45	33
AAP34180	167	94.7	9.00E-19	164	59	35	BAB18968	200	75.9	4.00E-13	134	46	33
P50612	494	94.4	1.00E-18	166	56	33	BAB18961	200	75.9	4.00E-13	134	45	33
P50612	494	61.2	1.00E-08	299	71	23	BAB18958	200	75.9	4.00E-13	134	45	33
CAA81035	134	93.8	2.00E-18	154	59	38	BAB18950	200	75.9	4.00E-13	134	45	33
NP_867573	685	92.8	3.00E-18	169	62	36	BAB18949	200	75.9	4.00E-13	134	45	33
NP_867573	685	85.5	5.00E-16	126	56	43	BAB18948	200	76.9	4.00E-13	134	46	33
CAC03723	503	92.4	4.00E-18	208	66	31	BAB18946	200	75.9	4.00E-13	134	45	33
CAC03723	503	66.2	3.00E-10	252	61	24	BAC16528	199	75.5	6.00E-13	133	45	33
NP_404370	401	92	6.00E-18	309	79	25	BAC16532	199	75.1	7.00E-13	133	45	33
ZP_00329910395	91.7	8.60E-18	285	73	25	BAC16525	199	75.1	7.00E-13	133	45	33	
ZP_00196929279	91.7	8.00E-18	288	79	27	BAC16348	200	74.7	1.00E-12	134	45	33	
NP_353572	306	91.3	1.00E-17	313	77	24	NP_907272	253	74.3	1.00E-12	278	65	23
NP_963427	293	91.3	1.00E-17	293	75	25	BAC76096	198	74.3	1.00E-12	133	44	33
ZP_0098298321	90.8	1.00E-17	302	85	28	BAC76095	198	74.3	1.00E-12	133	44	33	
AAA88923	501	98.5	2.00E-17	200	57	28	BAC16534	199	73.9	2.00E-12	133	44	33
AAA88923	501	56.8	3.00E-07	146	47	32	BAC16533	199	73.9	2.00E-12	133	44	33
BAC98372	302	90.5	2.00E-17	301	81	26	BAC16527	199	73.9	2.00E-12	133	44	33
AAC41457	482	90.5	2.00E-17	150	51	34	BAD18C56	199	73.9	2.00E-12	133	44	33
AAC41457	482	61.2	1.00E-08	299	71	23	BAD18C57	193	73.9	2.00E-12	133	44	33
NP_227847	258	89.7	3.00E-17	248	63	25	BAD18C55	199	73.9	2.00E-12	133	44	33
NP_554084	288	89.7	3.00E-17	290	75	25	AAP34191	159	73.2	3.00E-12	162	54	33
ZP_00269010110	89.7	5.00E-17	87	47	64	ZP_00270458269	72.8	4.00E-12	302	69	22		
NP_700301	282	88.6	6.00E-17	298	76	25	ZP_00289016275	72	6.00E-12	275	64	23	
AAA85132	102	88.6	6.00E-17	87	45	51	BAD38984	199	72	6.00E-12	133	43	32
AAV35032	531	87.4	1.00E-16	142	52	36	YP_162339	285	71.6	8.00E-12	282	56	19
AAV35032	531	63.9	2.00E-09	94	34	36	NP_353571	337	71.6	8.00E-12	324	75	23
NP_391420	298	86.7	2.00E-16	288	80	27	NP_531244	320	71.6	8.00E-12	324	75	23
NP_541127	282	86.7	2.00E-16	298	75	25	ZP_00183364320	71.6	8.00E-12	321	74	23	
ZP_00288130268	85.5	5.00E-16	295	76	25	CAF74789	295	71.6	8.00E-12	294	67	22	
NP_773505	313	85.1	7.00E-16	311	73	23	AAD41622	127	71.2	1.00E-11	100	41	41
NP_104150	328	84.7	9E-16	329	75	22	AAP34187	187	71.2	1.00E-11	160	53	33
BAC98373	302	84.3	1E-15	92	44	47	AAP34184	154	71.2	1.00E-11	160	53	33

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
AAP34192	155	69.7	3.00E-11	159	52	32	AAA86521	414	52	7.00E-06	108	32	29
CAF74790	263	69.7	3.00E-11	284	61	21	AAQ60546	309	58.9	5.00E-08	299	62	20
AAB48473	249	68.9	5.00E-11	288	68	23	ZP_00127207530	58.9	5.00E-08	135	40	29	29
AAC27443	249	68.9	5.00E-11	186	50	26	AAA23951	268	55.5	7.00E-08	46	29	60
AAC27443	249	38.5	7.60E-02	49	20	40	AAA23951	288	41.6	9.00E-03	307	79	25
AAA99141	214	68.9	5.00E-11	222	59	26	NP_772484	757	58.5	7.00E-08	273	72	26
NP_281892	249	68.6	7.00E-11	288	66	22	NP_772484	757	34.7	1.10E+00	109	29	26
NP_782319	316	68.2	9.00E-11	284	68	23	AAK58562	200	58.5	7.00E-08	206	50	24
ZP_00337274271	67.4	2.00E-10	284	64	22	BAA77313	179	58.5	7.00E-08	69	29	42	
AAP34189	154	67.4	2.00E-10	158	52	32	BAA77311	136	58.5	7.00E-08	69	29	42
ZP_00335243300	67	2.00E-10	302	74	24	NP_637307	401	58.2	9.00E-08	134	38	28	
AAM90629	117	67	2.00E-10	129	46	35	NP_637307	401	36.8	4.90E-01	119	31	26
ZP_00149794326	67	2.00E-10	322	67	20	AAU13838	410	58.2	9.00E-08	121	35	28	
NP_972952	415	66.6	3.00E-10	285	65	22	AAU13838	410	38.5	7.60E-02	260	55	21
CAA24655	38	66.2	3.00E-10	37	34	91	YP_083110	287	57.8	1.00E-07	242	57	23
NP_227898	304	65.1	8.00E-10	256	82	24	NP_978079	287	57.8	1.00E-07	241	60	24
AAM90636	126	65.1	8.00E-10	130	41	31	AAK58561	202	57.8	1.00E-07	208	51	24
AAM90630	125	65.1	8.00E-10	130	42	32	NP_772486	757	57.4	2.00E-07	255	72	28
AAQ75042	195	64.7	1.00E-09	202	52	25	NP_772486	757	34.3	1.40E+00	109	29	26
YP_013346	291	64.3	1.00E-09	288	61	21	AAS14965	190	57.4	2.00E-07	186	47	25
ZP_00049388242	63.9	2.00E-09	188	52	27	NP_212316	424	57.4	2.00E-07	132	38	28	
NP_464233	291	63.8	2.00E-09	288	61	21	NP_212316	424	35	8.40E-01	77	18	23
ZP_00229464291	63.5	2.00E-09	288	61	21	AAE81420	395	57.4	2.00E-07	138	38	27	
AAQ33575	375	63.5	2.00E-09	305	71	23	AAE81420	395	50.4	2.00E-05	149	43	28
AAC27442	254	63.5	2.00E-09	202	51	25	AAG14364	410	57.4	2.00E-07	418	87	20
AAC27442	254	38.5	7.60E-02	49	20	40	NP_384775	394	57	2.00E-07	138	38	27
AAC85633	416	63.2	3.00E-09	268	60	22	NP_384775	394	50.4	2.00E-05	87	29	33
AAQ75043	195	62.8	4.00E-09	202	51	25	AAS91573	80	57	2.00E-07	60	27	45
NP_384778	321	62.4	5.00E-09	321	68	21	AAF34186	144	57	2.00E-07	149	47	31
NP_642302	401	62.4	5.00E-09	134	40	29	AAF34181	139	57	2.00E-07	135	41	30
NP_642302	401	40.8	1.50E-02	118	31	26	AAU07039	424	56.6	3.00E-07	157	46	29
AAC33575	375	62.4	5.00E-09	305	71	23	AAU07039	424	35	8.40E-01	77	18	23
CAC03720	316	62.4	5.00E-09	141	46	32	AAP08515	287	56.6	3.00E-07	243	57	23
CAD11201	281	62.4	5.00E-09	108	38	35	ZP_00128974504	55.6	3.00E-07	113	32	28	28
CAD11198	281	62.4	5.00E-09	108	38	35	ZP_00128974504	43.5	2.00E-03	238	50	21	21
AAQ75041	195	62.4	5.00E-09	202	50	24	A32808	395	56.2	4.00E-07	138	37	26
NP_470057	291	62	6.00E-09	288	60	20	A32808	395	50.4	2.00E-05	149	43	28
AAB81422	321	62	6.00E-09	321	65	20	NP_521913	316	56.2	4.00E-07	305	55	21
AAM75948	462	61.6	8.00E-09	231	55	23	NP_746494	521	56.2	4.00E-07	130	41	31
AAQ75049	195	61.6	8.00E-09	202	50	24	NP_746494	521	41.6	9.00E-03	149	37	24
AAQ75048	195	61.6	8.00E-09	202	50	24	AAL98374	205	56.2	4.00E-07	157	42	26
AAQ75047	195	61.6	8.00E-09	202	50	24	ZP_00173445304	66.2	4.00E-07	301	68	22	22
AAQ75044	195	61.6	8.00E-09	202	50	24	AAU13837	400	55.8	5.00E-07	115	34	29
AAQ75040	195	61.6	8.00E-09	202	50	24	AAU13837	400	32.7	4.20E+00	254	52	20
BAA20927	195	61.6	8.00E-09	202	50	24	AAK15325	433	55.8	5.00E-07	132	36	27
AAK58558	202	61.2	1.00E-08	208	52	25	AAK15325	433	36.6	2.90E-01	167	45	26
ZP_00237160267	60.8	1.00E-08	243	62	25	AAA65584	64	55.8	5.00E-07	63	27	42	
AAK58560	202	60.8	1.00E-08	208	51	24	ZP_00278994407	55.5	6.00E-07	250	55	22	22
BAA20929	195	60.8	1.00E-08	202	50	24	NP_718794	403	56.1	8.00E-07	136	36	26
AAU13838	416	60.5	2.00E-08	124	36	29	NP_718794	403	34.3	1.40E+00	89	26	29
AAU13836	416	36.6	7.60E-02	260	55	21	NP_791768	530	55.1	8.00E-07	130	36	27
AAK58557	202	60.5	2.00E-08	208	50	24	NP_772485	757	55.1	8.00E-07	250	63	25
AAK58548	202	60.5	2.00E-08	208	50	24	NP_772485	757	33.9	1.90E+00	109	29	26
NP_348828	425	60.1	2.00E-08	289	65	22	CAB64773	399	55.1	8.00E-07	193	48	24
NP_346828	425	42.4	6.00E-03	310	69	22	CAB64773	399	39.7	3.40E-02	72	22	30
YP_126585	411	60.1	2.00E-08	147	39	26	BAA77309	94	55.1	8.00E-07	49	23	46
YP_095257	411	60.1	2.00E-08	147	39	26	NP_880130	510	54.7	1.00E-06	136	43	31
AAM90638	126	60.1	2.00E-08	124	41	33	YP_003363	422	54.7	1.00E-06	110	34	30
AAK58547	202	60.1	2.00E-08	208	49	23	NP_992941	337	54.3	1.00E-06	310	73	23
AAK58550	202	60.1	2.00E-08	208	49	23	NP_883789	510	54.3	1.00E-06	136	42	30
NP_104151	356	59.7	3.00E-08	357	81	22	NP_405374	326	54.3	1.00E-06	310	73	23
AAL99376	207	59.7	3.00E-08	162	42	25	ZP_00212898398	54.3	1.00E-06	111	29	26	26
AAM90534	126	59.7	3.00E-08	138	44	31	NP_799784	289	53.9	2.00E-06	258	55	21
YP_123558	411	59.3	4.00E-08	147	38	25	NP_772483	763	53.9	2.00E-06	186	47	25
YP_123558	411	32	7.10E+00	83	25	NP_772483	763	36.2	3.80E-01	109	29	26	
BAC75563	530	59.3	4.00E-08	130	38	29	NP_244487	395	53.9	2.00E-06	126	33	26
AAK58545	202	59.3	4.00E-08	208	49	23	CAA75368	168	53.9	2.00E-06	159	39	24
AAA86521	414	58.9	5.00E-08	270	64	23	BAU77319	121	53.9	2.00E-06	92	32	34

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	
BAA77315	123	53.9	2.00E-06	92	32	34	YP_04084910746	42.4	5.00E-03	340	71	20		
NP_353693	436	63.5	2.00E-06	256	59	23	YP_04084910746	36.6	2.90E-01	297	58	19		
NP_753263	317	53.5	2.00E-06	299	67	22	YP_04084910746	35.8	4.90E-01	301	63	20		
NP_531268	430	53.5	2.00E-06	256	59	23	YP_04084910746	33.5	2.40E+05	279	55	19		
AAB81421	396	63.5	2.00E-06	232	54	23	YP_04084910746	33.1	3.20E+00	317	68	21		
AAB81421	395	40	2.00E-02	87	24	27	YP_04084910746	31.6	9.30E+00	297	57	19		
BAA77301	123	53.5	2.00E-06	108	33	30	AAB81423	395	48.5	7.00E-05	285	66	23	
ZP_00317213517	53.1	3.00E-06	100	31	31		AAB81423	398	45.1	8.00E-04	146	38	26	
ZP_00317213517	40	2.60E-02	91	24	26	IORY	55	48.5	7.00E-05	50	24	48		
ZP_30273669412	52.8	4.00E-06	311	87	21	ZP_00167907404	48.5	7.00E-05	121	37	30			
ZP_30273669412	35.8	4.90E-01	146	35	23	ZP_00167907404	39.4	6.40E-01	98	23	23			
ZP_00055435521	52.8	4.00E-06	185	40	21	AAB89136	37	48.5	7.00E-05	38	23	63		
NP_384777	401	52.8	4.00E-06	189	48	25	AAG14365	411	48.1	1.00E-04	207	51	24	
NP_384777	401	40	2.60E-02	87	24	27	AAG14365	411	42.4	5.00E-03	116	27	23	
AAQ63645	159	52.4	5.00E-06	150	40	26	NP_384776	394	47.8	1.00E-04	137	34	24	
NP_415601	317	52	7.00E-06	305	70	22	NP_384776	394	49.8	5.00E-04	246	56	22	
AAQ62964	145	52	7.00E-06	144	38	26	AAF34780	3381	47.8	1.00E-04	286	55	19	
NP_421770	424	52	7.00E-06	164	35	21	AAF34780	3381	45.1	8.00E-04	295	58	19	
CAA75363	168	52	7.00E-06	189	38	23	AAF34780	3381	44.7	1.00E-03	274	50	18	
AAAG55829	317	51.6	9.00E-06	289	66	22	AAF34780	3381	42.7	4.00E-03	285	51	17	
NP_967522	361	51.6	9.00E-06	178	44	24	AAF34780	3381	41.2	1.20E-02	300	56	18	
YP_066426	757	51.6	9.00E-06	130	40	30	AAF34780	3381	39.7	3.40E-02	278	48	17	
AAM27194	181	51.6	9.00E-06	181	44	24	BAA05156	24	47.8	1.00E-04	24	24	100	
BAA77303	156	51.6	9.00E-06	88	27	NP_562800	451	47.4	2.00E-04	220	53	24		
B32808	395	51.2	1.00E-05	285	66	23	NP_455677	317	47.4	2.00E-04	140	38	27	
B32808	398	42	7.00E-03	87	23	NP_646141	9904	47	2.00E-04	306	64	20		
YP_111446	2634	51.2	1.00E-05	287	60	20	NP_646141	9904	43.1	3.00E-03	153	38	23	
YP_111446	2634	50.8	1.00E-05	281	59	20	NP_646141	9904	39.7	3.40E-02	332	71	21	
YP_111446	2634	47	2.00E-04	298	63	21	NP_646141	9904	38.3	4.40E-02	263	66	23	
YP_111446	2634	46.2	4.00E-04	291	60	20	NP_646141	9904	38.1	9.90E-02	279	56	20	
YP_111446	2634	45.1	8.00E-04	292	58	19	NP_646141	9904	35.4	6.40E-01	317	66	20	
YP_111446	2634	43.5	2.00E-03	284	58	20	NP_646141	9904	33.9	1.90E+00	297	57	19	
YP_111446	2634	43.5	2.00E-03	283	58	20	AAQ73340	142	47	2.00E-04	132	36	27	
YP_111446	2634	43.5	2.00E-03	289	59	20	YP_012160	282	47	2.00E-04	223	53	23	
YP_111446	2634	43.1	3.00E-03	292	59	20	AAB21164	31	47	2.00E-04	31	24	77	
YP_111446	2634	42	7.00E-03	281	63	22	NP_249778	439	46.5	3.00E-04	134	34	25	
YP_111446	2634	40	2.60E-02	295	59	20	NP_249778	439	32	7.10E+00	257	50	19	
YP_111446	2634	40	2.60E-02	298	59	19	ZP_00138675439	48.6	3.00E-04	134	34	25		
YP_111446	2634	39.3	4.40E-02	302	59	19	ZP_00314296822	46.2	4.00E-04	257	54	21		
YP_111446	2634	36.9	5.80E-02	283	53	18	NP_223758	257	46.2	4.00E-04	149	40	26	
YP_111446	2634	37.4	1.70E-01	278	48	17	YP_108306	1606	46.2	4.00E-04	296	66	22	
YP_111446	2634	35.8	4.90E-01	235	50	21	YP_108306	1606	38.1	9.90E-02	281	57	20	
YP_105520	459	51.2	1.00E-05	248	57	22	YP_108306	1606	38.1	9.90E-02	228	54	23	
YP_105520	459	41.6	9.00E-03	293	60	20	YP_108306	1606	38.1	9.90E-02	309	60	19	
YP_027819	207	51.2	1.00E-05	115	30	26	YP_108306	1606	37.4	1.70E-01	266	58	22	
NP_865849	651	51.2	1.00E-05	129	34	26	AAK57535	140	46.2	4.00E-04	126	35	27	
NP_947642	935	50.8	1.00E-05	284	70	24	CAC84733	78	46.2	4.00E-04	71	25	35	
NP_947642	935	34.3	1.40E+00	93	22	NP_267773	1072	45.4	6.00E-04	287	53	18		
NP_949268	888	50.8	1.00E-05	284	69	24	AA869370	89	46.2	4.00E-04	77	21	27	
NP_949268	888	38.9	5.80E-02	305	63	20	ZP_00290372765	45.8	5.00E-04	260	56	20		
YP_009743	523	50.4	2.00E-05	123	32	26	CAA75373	158	45.8	5.00E-04	139	33	23	
YP_009743	523	33.9	1.90E+00	171	36	21	CAA75370	154	45.8	5.00E-04	139	33	23	
ZP_00129004282	50.4	2.00E-06	286	68	23	CAA75369	159	45.8	5.00E-04	139	33	23		
NP_282040	750	50.1	3.00E-05	129	36	27	NP_267773	1072	45.4	6.00E-04	287	53	18	
NP_282040	750	41.6	9.00E-03	229	48	20	NP_267773	1072	33.5	2.40E+00	91	21	23	
ZP_00242042398	48.7	3.00E-05	215	53	24	AAQ64681	134	45.4	6.00E-04	132	35	26		
ZP_00242042398	39.7	3.40E-02	140	35	25	NP_228520	407	45.4	6.00E-04	290	63	21		
AAS66690	430	48.3	4.00E-05	134	34	25	ZP_003230792334	45.4	6.00E-04	274	50	18		
AAS66690	430	48.5	7.00E-05	130	33	25	ZP_003230792334	38.7	3.40E-02	278	49	17		
BAA77305	78	48.9	6.00E-05	74	28	37	ZP_003230792334	39.3	4.40E-02	293	54	18		
ZP_00283356523	48.5	7.00E-05	162	43	26	ZP_003230792334	38.5	7.60E-02	294	53	18			
ZP_00283356523	35	8.40E-01	106	23	21	ZP_003230792334	37	2.20E-01	280	52	18			
NP_460165	317	48.5	7.00E-05	140	38	27	ZP_003230792334	34.7	1.10E+00	261	56	19		
YP_150904	317	48.5	7.00E-06	140	38	27	CAA75376	183	45.4	6.00E-04	139	33	23	
YP_128296	304	48.5	7.00E-05	293	53	18	CAA75374	155	45.4	6.00E-04	139	33	23	
YP_040849	10746	48.5	7.00E-05	303	63	20	CAA75372	159	45.4	6.00E-04	139	33	23	

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
CAA75367	154	45.4	6.00E-04	139	33	23	ZP_00326048509	43.1	3.00E-03	273	63	23	
CAA75365	157	45.4	6.00E-04	139	33	23	ZP_00326048509	32	7.10E+00	99	20	20	
CAA75364	159	45.4	6.00E-04	139	33	23	CAA36029	21	43.1	3.00E-03	21	21	100
AAD25047	156	45.4	5.00E-04	143	36	25	ZP_00051799172	43.1	3.00E-03	136	31	22	
BAA02072	24	45.1	8.00E-04	24	23	95	AAP48611	107	43.1	3.00E-03	91	27	29
YP_111449	1530	45.1	8.00E-04	295	64	21	NP_207286	295	43.1	3.00E-03	150	42	28
YP_111449	1530	44.7	1.00E-03	292	64	21	NP_936893	989	43.1	3.00E-03	289	63	21
YP_111449	1530	44.3	1.00E-03	278	69	21	AAK39533	107	43.1	3.00E-03	91	27	29
YP_111449	1530	42.7	4.00E-03	320	69	21	AAQ59385	302	42.7	4.00E-03	262	55	13
YP_111449	1530	41.6	9.00E-03	238	59	20	NP_267008	1063	42.7	4.00E-03	291	56	19
YP_111449	1530	41.2	1.20E-02	284	60	21	NP_267008	1053	37	2.20E-01	231	42	18
YP_111449	1530	40	2.60E-02	250	51	20	NP_267008	1053	36.6	2.90E-01	281	63	22
YP_111449	1530	37.7	1.30E-01	250	50	20	YP_042074	1351	42.7	4.00E-03	304	59	19
YP_111449	1530	37	2.20E-01	263	52	19	CAA61527	436	42.7	4.00E-03	99	27	27
YP_111449	1530	34.7	1.10E+00	257	56	21	AAK39530	107	42.7	4.00E-03	81	26	32
CAA75375	151	45.1	8.00E-04	139	33	23	AAB69368	89	42.7	4.00E-03	77	20	26
CAA61528	436	45.1	8.00E-04	106	29	27	NP_939989	1254	42.4	5.00E-03	169	38	22
AAF67137	159	45.1	8.00E-04	146	37	25	YP_105401	1535	42.4	5.00E-03	300	63	21
NP_949252	886	44.7	1.00E-03	194	49	25	YP_105401	1535	36.2	3.80E-01	307	64	20
NP_949252	886	33.9	1.90E+00	102	25	24	YP_105401	1535	36.8	4.90E-01	217	46	21
YP_155526	404	44.7	1.00E-03	130	35	25	YP_065260	667	42.4	6.00E-03	263	54	20
YP_155526	404	32	7.10E+00	196	41	20	NP_906736	846	42.4	5.00E-03	168	37	22
NP_929192	321	44.7	1.00E-03	318	74	23	ZP_001432362005	42.4	5.00E-03	256	59	23	
NP_647392	2275	44.7	1.00E-03	287	53	18	AAK39535	107	42.4	5.00E-03	81	25	30
NP_647392	2275	34.3	1.40E+00	216	42	19	AAK39534	107	42.4	5.00E-03	81	25	30
NP_647392	2275	34.3	1.40E+00	299	60	20	AAB69365	89	42.4	5.00E-03	77	20	25
NP_371958	6713	44.7	1.00E-03	303	61	20	NP_523179	1309	42	7.00E-03	264	60	22
NP_371958	6713	42	7.00E-03	153	36	23	NP_523179	1309	36.2	3.80E-01	260	56	21
NP_371958	6713	41.2	1.20E-02	332	72	21	NP_523179	1309	35.8	4.90E-01	247	57	23
NP_371958	6713	35.4	6.40E-01	276	53	19	AAF95046	672	42	7.00E-03	320	64	20
NP_371958	6713	33.1	3.20E+00	297	57	19	NP_488975	661	42	7.00E-03	268	63	23
AAAP48612	107	44.7	1.00E-03	94	24	25	NP_784110	1377	42	7.00E-03	285	64	22
NP_374548	6713	44.7	1.00E-03	303	61	20	AAL58470	2283	42	7.00E-03	239	55	18
NP_374548	6713	42	7.00E-03	153	36	23	AAL58470	2283	35.4	6.40E-01	289	58	21
NP_374548	6713	41.2	1.20E-02	332	72	21	AAL58470	2283	33.9	1.90E+00	285	52	18
NP_374548	6713	35.4	6.40E-01	276	53	19	AAL58470	2283	33.5	2.40E+00	270	44	16
NP_374548	6713	33.1	3.20E+00	297	57	19	AAQ97872	309	41.6	9.00E-03	305	71	23
YP_106908	410	44.7	1.00E-03	124	28	22	YP_129124	395	41.6	9.00E-03	119	25	21
YP_044654	2275	44.7	1.00E-03	287	53	18	CAC84729	68	41.6	9.00E-03	65	24	36
YP_044654	2275	34.3	1.40E+00	216	42	19	AAA27074	20	41.6	9.00E-03	20	20	100
AAF57139	159	44.7	1.00E-03	146	37	25	AAAB69357	89	41.6	9.00E-03	74	24	32
AAK39536	107	44.7	1.00E-03	94	24	25	ZP_00089080401	41.6	9.00E-03	122	36	29	
AAK39541	107	44.7	1.00E-03	94	24	25	ZP_00089080401	38.5	7.60E-02	139	36	25	
AAA27080	284	44.3	1.00E-03	30	21	70	NP_465811	1787	41.2	1.20E-02	268	58	21
AAA27080	284	37.4	1.70E-01	31	18	NP_002143701439	41.2	1.20E-02	297	72	24		
YP_110805	1653	44.3	1.00E-03	269	58	21	ZP_002143701439	32.3	5.40E+00	272	56	20	
YP_110805	1653	40	2.60E-02	254	56	22	AAAB69377	89	41.2	1.20E-02	74	23	31
YP_110805	1653	37	2.20E-01	287	58	20	AAP07456	893	40.8	1.50E-02	263	49	18
YP_110805	1653	36.2	3.80E-01	274	56	20	CAA90950	20	40.8	1.50E-02	20	20	100
YP_110805	1653	34.7	1.10E+00	265	52	19	AAS91595	44	40.8	1.50E-02	38	18	47
CAA75371	159	44.3	1.00E-03	139	33	23	ZP_00063096901	40.8	1.50E-02	241	60	24	
AAD21057	4545	44.3	1.00E-03	228	52	22	NP_251152	5627	40.4	2.00E-02	329	67	20
ZP_001627551140	44.3	1.00E-03	295	62	21	NP_251152	5627	34.3	1.40E+00	235	50	21	
T30822	1366	44.3	1.00E-03	284	53	18	NP_251152	5527	32.7	4.20E+00	312	60	19
AAF94608	4558	43.9	2.00E-03	228	52	22	AAM09955	3692	40.4	2.00E-02	278	64	23
CAA57229	962	43.9	2.00E-03	284	53	18	AAM09955	3692	39.3	4.40E-02	321	76	23
CAA57228	1344	43.9	2.00E-03	284	53	18	AAS91595	44	40.4	2.00E-02	38	18	47
CAA57228	1344	43.9	2.00E-03	284	53	18	AAB869376	89	40.4	2.00E-02	64	21	32
CAA57228	1344	43.5	2.00E-03	254	53	18	AAB869369	89	40.4	2.00E-02	64	21	32
CAA57228	1344	43.1	3.00E-03	269	50	18	AAB869354	89	40.4	2.00E-02	64	21	32
AAK39539	107	43.9	2.00E-03	94	24	25	ZP_00288079749	40	2.60E-02	235	51	21	
AAB69366	89	43.9	2.00E-03	77	21	27	AAA92491	20	40	2.60E-02	20	18	90
AAA81014	716	43.9	2.00E-03	284	53	18	YP_142319	769	40	2.60E-02	302	64	21
AAA81014	716	43.1	3.00E-03	269	50	18	YP_140402	710	40	2.60E-02	302	64	21
CAF74791	410	43.9	2.00E-03	224	46	20	NP_373178	2271	40	2.60E-02	287	52	18
AAAP48610	107	43.5	2.00E-03	91	28	30	NP_373178	2271	34.3	1.40E+00	216	42	19
AAAP48609	107	43.5	2.00E-03	91	28	30	NP_373178	2271	32	7.10E+00	303	62	20
AAB69367	89	43.5	2.00E-03	77	20	25	AAO06483	623	40	2.60E-02	114	31	27

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
YP_114536	437	40	2.50E-02	83	27	32	NP_419715	307	38.1	9.90E-02	73	21	28
YP_034700	953	40	2.60E-02	301	58	19	NP_207906	228	38.1	9.90E-02	110	32	29
NP_836493	623	40	2.60E-02	114	31	27	ZP_003235731130	38.1	9.90E-02	292	65	22	
NP_764683	9439	40	2.60E-02	130	27	20	ZP_003235731130	32.3	5.40E+00	281	57	20	
NP_764683	9439	38.5	7.60E-02	169	43	25	ZP_00317192676	37.7	1.30E-01	168	35	20	
NP_764683	9439	36.2	3.80E-01	265	55	20	CAA60556	61	37.7	1.30E-01	43	16	37
NP_764683	9439	36.2	3.80E-01	238	61	21	NP_771092	696	37.7	1.30E-01	134	39	29
NP_764683	9439	36.2	3.80E-01	255	48	18	NP_214145	422	37.7	1.30E-01	186	40	21
NP_764683	9439	35.4	6.40E-01	316	51	19	NP_765968	681	37.7	1.30E-01	176	45	25
NP_764683	9439	34.7	1.10E+00	302	60	19	ZP_003125671475	37.7	1.30E-01	273	58	21	
NP_764683	9439	33.9	1.90E+00	314	55	20	ZP_00240971564	37.7	1.30E-01	107	24	22	
NP_764683	9439	33.9	1.90E+00	146	40	27	ZP_001952635372	37.7	1.30E-01	183	41	22	
NP_764683	9439	33.1	3.20E+00	181	46	25	BAD13420	718	37.7	1.30E-01	89	18	20
NP_764683	9439	33.1	3.20E+00	151	38	25	AAB19816	20	37.7	1.30E-01	20	18	90
NP_764683	9439	32.3	5.40E+00	329	68	20	NP_949265	623	37.4	1.70E-01	277	53	19
NP_764683	9439	31.6	9.30E+00	292	53	18	ZP_00315446537	37.4	1.70E-01	263	54	20	
AAC08547	402	39.7	3.40E-02	148	34	22	ZP_0028945115245	37.4	1.70E-01	272	65	23	
ZP_001236916388	39.7	3.40E-02	352	74	21	ZP_00273576527	37.4	1.70E-01	90	25	27		
ZP_000642471003	39.7	3.40E-02	307	71	23	ZP_00273576527	33.9	1.90E+00	83	22	26		
AAC09051	130	39.7	3.40E-02	97	26	26	NP_800021	623	37.4	1.70E-01	117	31	26
AAC09050	130	39.7	3.40E-02	97	26	26	NP_772493	627	37.4	1.70E-01	229	49	21
AAC09048	130	39.7	3.40E-02	97	26	26	AAP78292	855	37.4	1.70E-01	138	32	23
AABG2741	425	39.7	3.40E-02	135	34	25	ZP_00281310507	37.4	1.70E-01	119	31	26	
AA862741	425	32	7.10E+00	78	18	ZP_00289133779	37	2.20E-01	198	38	19		
AAR21284	163	39.7	3.40E-02	97	26	ZP_00289133779	36.2	3.80E-01	89	26	29		
AAR21282	161	39.7	3.40E-02	97	26	NP_471790	927	37	2.20E-01	281	59	20	
ZP_00319871233	39.3	4.40E-02	139	37	26	NP_471790	927	31.6	9.30E+00	210	46	21	
NP_908061	118	39.3	4.40E-02	103	29	28	NP_522634	3322	37	2.20E-01	228	52	22
ZP_002335911787	39.3	4.40E-02	261	60	22	NP_404364	307	37	2.20E-01	253	52	20	
AAC09046	130	39.3	4.40E-02	97	26	26	NP_336479	342	37	2.20E-01	87	25	28
1QOY	318	39.3	4.40E-02	77	27	35	NP_248731	3535	37	2.20E-01	286	62	21
NP_798012	3240	38.9	5.80E-02	269	58	21	YP_086441	693	37	2.20E-01	120	30	25
YP_015833	662	38.9	5.80E-02	167	43	25	NP_757754	161	37	2.20E-01	77	23	29
ZP_002814314726	38.9	5.80E-02	310	62	20	NP_670743	368	37	2.20E-01	253	52	20	
ZP_002814314726	38.1	9.90E-02	308	61	19	NP_603291	1724	37	2.20E-01	131	34	25	
ZP_002814314726	37.4	1.70E-01	272	54	19	ZP_00299009010	37	2.20E-01	260	53	20		
ZP_002814314726	36.6	2.90E-01	275	56	20	ZP_00219646252	37	2.20E-01	74	21	28		
ZP_002814314726	36.6	2.90E-01	254	53	20	ZP_00063136721	37	2.20E-01	298	58	19		
ZP_002814314726	35.4	6.40E-01	279	55	19	NP_799785	346	36.6	2.90E-01	116	29	26	
ZP_002814314726	35	8.40E-01	319	59	18	NP_736300	290	36.6	2.90E-01	122	30	24	
ZP_002814314726	34.3	1.40E-00	324	59	18	AAP96352	1119	36.6	2.90E-01	143	35	24	
ZP_002114934727	38.9	5.80E-02	274	60	21	NP_359556	1902	36.6	2.90E-01	282	60	21	
ZP_002047941979	38.9	5.80E-02	235	47	20	NP_253231	1417	36.6	2.90E-01	201	49	24	
ZP_002047941979	34.7	1.10E+00	235	50	21	NP_932208	603	36.6	2.90E-01	53	17	32	
ZP_002047941979	33.1	3.20E+00	312	61	19	CAA60548	61	36.6	2.90E-01	43	15	34	
AAC43555	35	38.9	5.80E-02	34	21	AAT28336	160	36.6	2.90E-01	165	34	20	
ZP_00208729459	38.5	0.076	278	63	22	AAT28333	160	36.6	2.90E-01	166	34	20	
NP_928380	567	38.5	0.076	257	58	22	AAT28333	180	36.6	2.90E-01	143	35	24
YP_115646	760	38.5	7.60E-02	144	32	22	ZP_0021898486	36.6	2.90E-01	282	60	21	
YP_087940	5399	38.5	7.60E-02	312	72	23	ZP_00149774452	36.6	2.90E-01	265	56	21	
YP_087940	5399	37.4	1.70E-01	256	45	17	AAP34183	119	36.6	2.90E-01	115	31	26
YP_087940	5399	37.4	1.70E-01	256	45	17	ZP_00319246821	36.2	3.80E-01	65	19	29	
YP_087940	5399	36.2	3.80E-01	273	58	21	ZP_00284479501	36.2	3.80E-01	262	52	19	
YP_087940	5399	36.2	3.80E-01	273	58	21	ZP_00005278425	36.2	3.80E-01	193	50	25	
YP_087940	5399	35.8	4.90E-01	274	61	22	NP_522101	3552	36.2	3.80E-01	209	45	21
YP_087940	5399	35	8.40E-01	294	58	19	NP_964415	1096	36.2	3.80E-01	265	56	21
YP_087940	5399	33.9	1.90E+00	269	57	19	AAG31286	1363	36.2	3.80E-01	193	50	25
YP_087940	5399	33.5	2.40E+00	269	57	19	ZP_00313063880	36.2	3.80E-01	181	39	21	
NP_781518	518	38.5	7.60E-02	302	64	21	AAT28334	160	36.2	3.80E-01	166	34	20
AAM46179	146	38.5	7.60E-02	95	25	26	ZP_00182337425	36.2	3.80E-01	245	41	16	
AAM46178	147	38.5	7.60E-02	95	25	26	ZP_00160617717	36.2	3.80E-01	289	62	21	
CAC61118	1564	38.5	7.60E-02	275	60	21	ZP_00160617717	31.6	9.30E+00	248	46	18	
ZP_00213493557	38.5	7.60E-02	223	51	22	ZP_001380311417	36.2	3.80E-01	201	49	24		
ZP_00224552561	38.5	7.60E-02	223	53	23	AAR37720	394	36.2	3.80E-01	146	39	26	
ZP_00355727801	38.5	7.60E-02	317	71	22	NP_561526	1109	35.8	4.90E-01	287	53	18	
ZP_001404443443	38.5	7.60E-02	286	62	21	NP_561526	1109	32	7.10E+00	181	40	22	
AAC43560	36	38.5	7.60E-02	36	19	NP_349224	664	35.8	4.90E-01	272	59	21	
AAB69373	89	38.5	7.60E-02	74	22	AAP56614	499	35.8	4.90E-01	183	40	21	

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
YP_133518	665	35.8	4.90E-01	111	33	29	NP_695350	459	34.3	1.40E+00	271	54	19
YP_071943	3378	35.8	4.90E-01	289	61	21	NP_348729	570	34.3	1.40E+00	162	36	22
YP_049813	317	35.8	4.90E-01	266	61	22	YP_046173	335	34.3	1.40E+00	117	26	22
AAT28337	180	35.8	4.90E-01	166	34	20	NP_252997	632	34.3	1.40E+00	245	43	17
AAA27141	17	35.8	4.90E-01	17	17	100	YP_066477	393	34.3	1.40E+00	148	33	22
ZP_00270453727	35.4	6.40E-01	239	49	20	YP_064761	880	34.3	1.40E+00	126	30	23	
AAQ59977	458	35.4	6.40E-01	256	61	22	YP_064723	674	34.3	1.40E+00	248	48	19
NP_773497	491	35.4	6.40E-01	121	28	23	NP_935137	626	34.3	1.40E+00	299	52	17
NP_267038	799	35.4	6.40E-01	301	71	23	NP_866060	3056	34.3	1.40E+00	120	34	28
NP_478503	1487	35.4	6.40E-01	249	49	19	NP_866050	3056	33.1	3.20E+00	116	34	29
YP_086339	564	35.4	6.40E-01	107	23	21	ZP_003233981656	34.3	1.4		187	41	21
NP_207948	1230	35.4	6.40E-01	292	62	21	ZP_00308827366	34.3	1.40E+00	138	35	25	
YP_039058	564	35.4	6.40E-01	107	23	21	ZP_00137780616	34.3	1.40E+00	245	43	17	
NP_864533	7718	35.4	6.40E-01	240	52	21	ZP_00124017629	34.3	1.40E+00	254	46	18	
NP_765804	2310	35.4	6.40E-01	290	52	17	ZP_00110825475	34.3	1.40E+00	140	35	25	
NP_765787	676	35.4	6.40E-01	117	33	28	NP_00046942967	33.9	1.90E+00	134	35	26	
ZP_00323209769	35.4	6.40E-01	148	35	23	YP_080405	650	33.9	1.90E+00	87	22	25	
ZP_00288038697	35.4	6.40E-01	294	56	19	NP_470074	599	33.9	1.90E+00	288	59	20	
CAE46764	510	35.4	6.40E-01	95	29	30	NP_927899	4582	33.9	1.90E+00	257	61	23
CAE46762	506	35.4	6.40E-01	95	29	30	NP_815453	533	33.9	1.90E+00	125	35	28
T08513	619	35.4	6.40E-01	173	37	21	NP_562046	933	33.9	1.90E+00	259	51	19
AAP34195	125	35.4	6.40E-01	129	35	27	NP_267826	901	33.9	1.90E+00	281	56	19
ZP_00338261589	35	8.40E-01	62	20	32	YP_126425	657	33.9	1.90E+00	121	29	23	
NP_718838	667	35	8.40E-01	98	20	20	YP_131301	638	33.9	1.90E+00	73	16	21
YP_128297	361	35	8.40E-01	358	68	19	YP_030040	660	33.9	1.90E+00	105	25	23
YP_132326	529	35	8.40E-01	332	69	20	NP_981473	584	33.9	1.90E+00	109	24	22
NP_812240	868	35	8.40E-01	150	44	29	NP_784971	983	33.9	1.9	299	53	19
NP_964984	4734	35	8.40E-01	316	65	20	NP_783077	417	33.9	1.90E+00	103	24	23
NP_964984	4734	33.1	3.20E+00	153	40	26	NP_965011	912	33.9	1.90E+00	214	49	22
AAN33150	434	35	8.40E-01	127	36	27	NP_864462	1000	33.9	1.90E+00	125	28	22
ZP_00144937646	35	8.40E-01	243	44	18	AAP02960	599	33.9	1.90E+00	119	30	25	
ZP_00222048485	35	8.40E-01	88	23	26	BAB20920	1172	33.9	1.90E+00	283	55	19	
ZP_00220113801	35	8.40E-01	88	23	26	ZP_002991151200	33.5	2.4		61	17	27	
ZP_00137782629	35	8.40E-01	245	43	17	ZP_00056271661	33.5	2.40E+00	278	54	19		
ZF_00121836231	35	8.40E-01	165	39	23	AAG56007	973	33.5	2.40E+00	185	42	22	
AAQ19127	2343	35	8.40E-01	239	49	20	AAF81209	277	33.5	2.40E+00	199	45	22
NP_948001	587	34.7	1.10E+00	121	36	29	NP_794375	539	33.5	2.40E+00	128	23	17
ZP_00319892754	34.7	1.10E+00	284	62	21	YP_115527	534	33.5	2.40E+00	80	24	30	
ZP_00319892754	33.1	3.20E+00	261	51	19	AAK93934	736	33.5	2.40E+00	89	20	22	
ZP_00311897602	34.7	1.10E+00	269	56	20	NP_309677	971	33.5	2.4	185	42	22	
YP_160437	862	34.7	1.10E+00	68	24	35	NP_936396	542	33.5	2.40E+00	300	62	20
ZP_00336629824	34.7	1.10E+00	149	35	23	NP_637327	396	33.5	2.40E+00	123	21	17	
NP_802026	593	34.7	1.10E+00	118	33	27	NP_984159	982	33.5	2.40E+00	203	47	23
NP_924800	1241	34.7	1.10E+00	73	24	32	ZP_00217082977	33.5	2.40E+00	109	32	29	
NP_463823	500	34.7	1.10E+00	233	47	20	ZP_00106687978	33.5	2.40E+00	112	29	25	
NP_645581	946	34.7	1.10E+00	219	44	20	AAB21165	20	33.5	2.40E+00	20	16	80
NP_792425	633	34.7	1.10E+00	283	50	17	BAA23410	632	33.5	2.40E+00	229	43	18
NP_561978	344	34.7	1.10E+00	232	49	21	ZP_00290912684	33.1	3.20E+00	130	24	18	
NP_350172	783	34.7	1.10E+00	129	29	ZP_00290912684	32.3	5.40E+00	188	36	18		
NP_349191	722	34.7	1.10E+00	143	36	25	YP_147975	435	33.1	3.20E+00	121	32	26
NP_349191	722	33.1	3.20E+00	292	57	19	ZP_00338277837	33.1	3.20E+00	109	26	23	
AAO10484	626	34.7	1.10E+00	299	52	17	AAU01878	1363	33.1	3.20E+00	175	40	22
NP_642322	396	34.7	1.10E+00	123	22	NP_798538	678	33.1	3.20E+00	127	27	21	
YP_133410	577	34.7	1.10E+00	317	61	19	NP_790701	498	33.1	3.20E+00	162	41	25
NP_269527	594	34.7	1.10E+00	118	33	27	NP_768559	816	33.1	3.20E+00	62	19	30
YP_060520	893	34.7	1.10E+00	118	33	27	NP_415890	1122	33.1	3.20E+00	147	37	25
YP_042878	928	34.7	1.10E+00	219	44	20	AAO08482	622	33.1	3.20E+00	120	30	25
YP_012922	500	34.7	1.10E+00	233	47	20	AAO10818	675	33.1	3.20E+00	125	29	23
NP_051325	222	34.7	1.10E+00	114	30	26	NP_889112	532	33.1	3.20E+00	250	48	19
NP_782459	569	34.7	1.10E+00	305	61	20	YP_129467	640	33.1	3.20E+00	137	31	22
NP_607632	594	34.7	1.10E+00	118	33	27	YP_133516	467	33.1	3.20E+00	127	26	20
ZP_00233976541	34.7	1.10E+00	233	47	20	YP_132040	542	33.1	3.20E+00	107	27	25	
ZP_00212291306	34.7	1.10E+00	130	38	29	YP_076807	558	33.1	3.20E+00	118	30	25	
AAK84427	542	34.7	1.10E+00	233	47	20	NP_253000	629	33.1	3.20E+00	245	42	17
AAA87447	232	34.7	1.10E+00	149	33	22	YP_063897	576	33.1	3.20E+00	263	46	17
YP_162386	564	34.3	1.40E+00	242	55	22	YP_041192	424	33.1	3.20E+00	147	36	24
NP_469464	1788	34.3	1.40E+00	290	65	22	NP_936490	522	33.1	3.20E+00	120	30	25
NP_792202	620	34.3	1.40E+00	128	31	24	NP_786249	1106	33.1	3.20E+00	330	64	19

Fig. 25B-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
NP_777771	638	33.1	3.20E+00	121	29	23	YP_032604	1872	32.3	5.40E+00	118	31	26
NP_764984	3692	33.1	3.2	148	30	20	NP_782523	471	32.3	5.40E+00	107	20	18
NP_764984	3692	32.3	5.40E+00	264	62	19	NP_765908	300	32.3	5.40E+00	132	29	21
NP_600104	441	33.1	3.20E+00	280	63	22	NP_765785	952	32.3	5.40E+00	195	37	18
NP_693909	466	33.1	3.20E+00	126	24	19	NP_606940	628	32.3	5.40E+00	271	60	22
NP_808047	400	33.1	3.20E+00	108	29	26	CAF32691	1828	32.3	5.40E+00	228	56	24
NP_604026	1193	33.1	3.20E+00	253	52	20	CAD6598	1184	32.3	5.40E+00	270	49	18
NP_541885	131	33.1	3.20E+00	62	24	38	ZP_002394591038	32.3	5.40E+00	243	52	21	
gi 42627767 pe CAD89361.1 TPA: SbcC exonuclease [Bradyrhizobium japonicum]	1261	33.1	3.20E+00	62	33.1	3.20E+00	ZP_00238267289	32.3	5.40E+00	156	31	19	
	19	30					ZP_002326811066	32.3	5.40E+00	277	57	20	
CAA64859	624	33.1	3.20E+00	284	47	16	CAB99193	326	32.3	5.40E+00	104	27	25
CAA81206	955	33.1	3.20E+00	239	47	19	AAT41983	427	32.3	5.40E+00	245	48	19
BAB98259	413	33.1	3.20E+00	280	63	22	AAF25839	177	32.3	5.40E+00	86	23	26
AAD43469	912	33.1	3.20E+00	240	48	20	AAS93940	693	32.3	5.40E+00	137	29	21
ZP_00137783633	33.1	3.20E+00	245	42	17	ZP_002188313513	32.3	5.40E+00	204	51	25		
ZP_00123950634	33.1	3.20E+00	96	24	25	ZP_00171926505	32.3	5.40E+00	253	48	18		
ZP_00128148551	33.1	3.20E+00	256	46	17	ZP_00201727822	32.3	5.40E+00	306	59	19		
ZP_003450801463	33.1	3.20E+00	274	69	25	P55116	953	32.3	5.40E+00	233	48	20	
AAG60897	896	33.1	3.20E+00	62	19	30	ZP_00149920505	32.3	5.40E+00	139	31	22	
PC6003	624	33.1	3.2	284	47	16	ZP_00133930158	32.3	5.40E+00	106	26	24	
P76072	1120	33.1	3.20E+00	147	37	25	ZP_00098108451	32.3	5.40E+00	139	33	23	
BAA23413	629	33.1	3.20E+00	245	42	17	AAK27341	3241	32.3	5.40E+00	102	24	23
ZP_00326431563	32.7	4.20E+00	307	70	22	ZP_00312854422	32	7.10E+00	142	36	25		
CAA79304	933	32.7	4.20E+00	145	30	20	ZP_00298543733	32	7.10E+00	278	50	17	
NP_252999	629	32.7	4.20E+00	127	32	25	ZP_003410411311	32	7.10E+00	230	37	16	
AAM01202	183	32.7	4.20E+00	121	29	23	ZP_00208481932	32	7.10E+00	211	43	20	
ZP_000531851136	32.7	4.20E+00	223	51	22	ZP_00046132979	32	7.10E+00	143	32	22		
NP_993634	2578	32.7	4.20E+00	290	61	21	NP_464751	1066	32	7.10E+00	277	55	20
YP_081346	561	32.7	4.20E+00	99	20	20	NP_BB5521	1195	32	7.10E+00	65	18	27
NP_471252	1186	32.7	4.20E+00	315	63	20	AAQ60245	251	32	7.10E+00	98	30	30
NP_469482	290	32.7	4.20E+00	82	21	35	NP_768828	432	32	7.10E+00	164	35	21
NP_464250	601	32.7	4.20E+00	193	39	20	NP_699533	317	32	7.10E+00	56	22	39
NP_929013	190	32.7	4.20E+00	133	34	25	NP_348690	371	32	7.1	147	35	23
NP_793014	6274	32.7	4.2	246	51	20	NP_207534	493	32	7.1	122	32	26
NP_793015	541	32.7	4.20E+00	93	28	30	NP_249777	683	32	7.1	303	72	23
NP_769573	564	32.7	4.20E+00	157	36	22	YP_060448	1039	32	7.1	193	42	21
NP_563507	721	32.7	4.20E+00	112	26	22	YP_041227	2189	32	7.10E+00	149	35	23
NP_406024	2635	32.7	4.20E+00	290	61	21	YP_010811	580	32	7.10E+00	206	44	21
NP_939703	888	32.7	4.20E+00	123	31	25	NP_784951	3360	32	7.10E+00	76	21	27
AAP56776	742	32.7	4.20E+00	186	40	21	NP_765498	495	32	7.10E+00	74	16	21
NP_882071	1175	32.7	4.20E+00	65	18	27	NP_757771	1378	32	7.10E+00	73	22	30
NP_419323	622	32.7	4.20E+00	235	47	20	NP_688852	443	32	7.10E+00	66	17	25
YP_108428	634	32.7	4.20E+00	266	52	19	AAO19442	5431	32	7.10E+00	206	44	21
NP_109825	228	32.7	4.20E+00	154	29	18	ZP_002842243286	32	7.10E+00	283	62	21	
YP_013363	601	32.7	4.20E+00	193	39	20	CAA05367	577	32	7.10E+00	163	32	19
NP_669014	2579	32.7	4.20E+00	290	61	21	BAD51767	1173	32	7.10E+00	283	53	18
ZP_00284267481	32.7	4.20E+00	77	21	27	BAC57543	289	32	7.1	135	33	24	
ZP_00280424428	32.7	4.20E+00	196	50	25	ZP_00355478659	32	7.1	128	32	25		
AAM01201	183	32.7	4.20E+00	121	29	ZP_00215525494	32	7.1	118	25			
BAA23412	629	32.7	4.20E+00	127	32	ZP_00215989433	32	7.1	61	20			
NP_326456	750	32.3	5.40E+00	121	27	ZP_00211381462	32	7.1	89	23			
CAA55199	545	32.3	5.40E+00	238	47	ZP_00202082371	32	7.1	115	31			
NP_391003	662	32.3	5.40E+00	322	80	18	ZP_00154800558	32	7.1	139	31		
A42292	818	32.3	5.40E+00	103	29	ZP_00348485544	32	7.1	93	22			
BAA02196	777	32.3	5.40E+00	134	35	ZP_00138674683	32	7.1	303	72			
AAP08880	1658	32.3	5.40E+00	149	33	ZP_00274787513	31.6	9.3	247	48			
AAP08880	1658	32.3	5.40E+00	139	27	NP_326624	1125	31.6	9.3	174	37		
NP_646487	424	32.3	5.40E+00	147	36	CAA41384	1134	31.6	9.3	147	31		
NP_718941	169	32.3	5.40E+00	160	36	AAD02406	269	31.6	9.3	81	21		
NP_716901	706	32.3	5.40E+00	273	59	ZP_00054847736	31.6	9.3	272	53			
NP_807002	542	32.3	5.40E+00	304	58	NP_941106	1062	31.6	9.3	221	48		
AAQ80540	2373	32.3	5.40E+00	255	55	NP_813981	522	31.6	9.3	137	33		
AAQ85839	690	32.3	5.40E+00	110	25	NP_794891	647	31.6	9.3	119	27		
NP_770343	582	32.3	5.4	270	50	NP_766736	691	31.6	9.3	101	30		
AAF96820	652	32.3	5.40E+00	120	31	NP_562167	327	31.6	9.3	253	47		
AAF94794	596	32.3	5.40E+00	80	19	NP_561782	2104	31.6	9.3	126	23		
NP_404241	3295	32.3	5.40E+00	289	61	NP_358633	1091	31.6	9.3	137	31		
						NP_349987	570	31.6	9.3	300	64	21	

Fig. 25C

Hit_ID	Length	Bitscore	Expected	Length	#ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#ident	%S
NP_929221_355	364	1.00E-100	356	207	58		NP_642301_399	195	2.00E-49	212	113	53	
ZP_00169997361	343	1.00E-93	361	201	55		NP_642301_399	103	1.00E-21	307	87	28	
NP_992909_369	318	4.00E-86	368	202	54		ZP_00236470278	196	2.00E-49	293	117	39	
NP_405406_369	318	4.00E-86	368	202	54		ZP_00236469286	195	3.00E-49	292	118	40	
NP_288384_585	305	4.00E-82	274	169	61		ZP_00244371402	195	4.00E-49	400	144	36	
NP_288384_585	142	4.00E-33	115	74	64		ZP_0031275272	193	1.00E-48	292	118	40	
NP_310689_585	305	4.00E-82	274	169	61		ZP_00335280509	193	1.00E-48	290	120	41	
NP_310689_585	142	4.00E-33	115	74	64		ZP_00335280509	114	1.00E-24	159	67	42	
NP_754230_595	298	3.00E-80	223	157	70		NP_637306_399	193	1.00E-48	241	119	49	
NP_754230_595	138	5.00E-32	265	106	41		NP_637306_399	104	1.00E-21	319	93	29	
NP_707809_550	293	9.00E-79	202	154	76		ZP_00244372404	193	1.00E-48	404	150	37	
NP_707809_550	140	9.00E-33	116	71	61		YP_065397_1128	192	3.00E-48	252	120	47	
NP_416433_498	287	8.00E-77	177	146	52		YP_065397_1128	107	2.00E-22	254	79	31	
NP_416433_498	149	3.00E-35	151	80	52		YP_001491_285	192	4.00E-48	302	112	37	
NP_841632_275	285	4.00E-76	297	159	53		NP_712598_285	192	4.00E-48	302	112	37	
NP_456520_505	284	7.00E-76	175	150	85		NP_231219_379	189	2.00E-47	375	130	34	
NP_456520_506	149	3.00E-35	91	76	83		YP_109915_388	189	2.00E-47	385	136	35	
NP_461898_508	283	9.00E-76	232	150	88		NP_831434_272	189	2.00E-47	298	114	38	
NP_461898_508	155	4.00E-37	100	81	81		YP_076801_275	189	3.00E-47	296	114	38	
NP_460912_495	280	8.00E-75	175	143	84		NP_967579_277	188	4.00E-47	301	117	38	
NP_460912_495	152	3.00E-36	98	79	80		YP_027349_287	188	5.00E-47	292	112	38	
NP_883763_392	272	2.00E-72	392	180	45		NP_672081_285	187	9.00E-47	300	113	37	
NP_879790_391	268	5.00E-71	391	182	46		ZP_00236468284	187	1.00E-46	295	111	37	
NP_889978_391	266	1.00E-70	391	183	45		ZP_00288129272	186	2.00E-46	296	112	37	
ZP_00091764573	254	8.00E-67	262	144	54		NP_967577_274	186	3.00E-46	292	115	39	
ZP_00091764573	139	4.00E-32	90	71	78		ZP_0020889296	184	6.00E-46	300	110	36	
ZP_00173465285	243	1.00E-65	297	145	48		YP_129125_382	184	7.00E-48	382	130	34	
NP_903549_282	248	6.00E-65	297	137	46		NP_219229_286	183	2.00E-46	298	114	38	
ZP_00288126272	247	7.00E-65	296	145	48		NP_348182_278	182	3.00E-45	292	115	39	
NP_303548_282	246	2.00E-64	297	137	46		YP_010650_298	182	3.00E-45	311	119	38	
ZP_00289013269	238	4.00E-62	292	133	45		NP_231773_376	182	4.00E-45	375	133	35	
ZP_00149772273	236	1.00E-61	296	131	44		ZP_00130180299	181	5.00E-45	295	107	36	
ZP_00289014271	236	2.00E-61	296	133	44		NP_219303_286	181	8.00E-45	304	107	35	
NP_718792_272	236	2.00E-61	297	140	47		NP_219305_285	181	8.00E-45	301	111	36	
ZP_00274388386	236	2.00E-61	157	122	73		NP_831435_273	180	1.00E-44	298	115	38	
ZP_00274388386	135	5.00E-31	92	68	73		YP_123618_475	179	2.00E-44	309	124	40	
ZP_00289011271	235	3.00E-61	296	133	44		YP_123618_475	110	1.00E-23	200	77	38	
ZP_00288132271	235	4.00E-61	295	136	46		YP_035879_367	179	2.00E-44	365	125	34	
ZP_00288131271	234	5.00E-61	295	136	46		YP_095369_475	178	4.00E-44	309	125	40	
ZP_00299645314	234	6.00E-61	296	137	46		YP_095369_475	109	2.00E-23	198	78	39	
ZP_00288133271	233	1.00E-60	295	135	45		YP_001839_281	177	9.00E-44	300	107	35	
NP_718793_273	233	1.00E-60	297	138	45		NP_971613_286	177	9.00E-44	304	108	35	
ZP_00288136271	230	9.00E-80	296	132	44		NP_902165_285	177	1.00E-43	299	104	34	
ZP_00211492272	229	2.00E-59	291	124	46		YP_086429_857	177	1.00E-43	231	106	46	
ZP_00289022272	228	3.00E-59	296	136	45		YP_066429_857	115	4.00E-25	227	81	35	
NP_954080_276	228	5.00E-59	298	133	44		YP_011656_297	177	1.00E-43	310	113	36	
NP_791772_282	223	1.00E-57	297	129	43		YP_126643_475	178	2.00E-43	309	122	39	
NP_622175_276	220	1.00E-56	296	134	45		YP_126643_475	110	1.00E-23	251	88	36	
ZP_00302655327	219	2.00E-56	329	133	40		YP_001450_282	176	2.00E-43	300	107	35	
NP_871067_421	219	2.00E-56	181	109	60		NP_970090_277	176	2.00E-43	295	110	37	
NP_871067_421	120	1.00E-26	346	109	31		NP_231775_378	176	2.00E-43	377	134	35	
ZP_00127282271	215	4.00E-56	283	128	45		YP_072597_336	176	3.00E-43	342	113	33	
YP_080862_310	216	4.00E-55	313	135	43		NP_212281_336	176	3.00E-43	342	113	33	
NP_244483_272	214	9.00E-55	294	124	42		NP_712599_282	176	3.00E-43	300	107	35	
ZP_00263336255	213	2.00E-54	269	125	46		NP_521943_273	175	3.00E-43	297	109	36	
ZP_00183380275	210	1.00E-33	299	128	42		NP_231818_377	175	3.00E-43	377	129	34	
NP_782313_275	209	2.00E-53	296	125	42		YP_013330_287	173	1.00E-42	296	107	36	
NP_693649_338	208	2.00E-53	338	131	38		NP_470041_287	173	1.00E-42	296	107	36	
ZP_00289017265	209	2.00E-53	285	125	43		NP_464217_287	173	1.00E-42	296	107	36	
ZP_00268012373	209	3.00E-53	282	117	41		NP_797167_384	173	1.00E-42	384	125	32	
NP_391416_204	208	5.00E-53	306	130	42		NP_535284_376	173	1.00E-42	375	121	32	
NP_301058_284	207	8.00E-53	301	119	39		NP_969824_277	172	3.00E-42	295	109	36	
NP_348820_275	207	1.00E-52	295	122	41		NP_972083_286	172	4.00E-42	304	106	34	
NP_798640_377	202	2.00E-51	376	132	36		NP_760797_377	171	7.00E-42	376	123	32	
ZP_00313276273	201	8.00E-51	295	120	40		ZP_00346385295	171	7.00E-42	292	98	33	
ZP_00236467307	200	1.00E-50	309	119	38		NP_348785_283	171	9.00E-42	290	104	35	
NP_967404_282	199	3.00E-50	295	124	42		NP_782278_280	170	1.00E-41	294	105	35	
NP_967404_277	196	2.00E-49	292	120	41		ZP_00279040381	170	1.00E-41	378	127	33	

Fig. 25C-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
NP_231774	377	169	2.00E-41	379	127	33	NP_282485	572	106	3.00E-22	171	63	36
YP_011296	297	169	2.00E-41	295	101	34	NP_282485	572	79.3	3.00E-14	132	48	36
NP_902681	372	167	7.00E-41	369	122	33	AAP78393	514	104	8.00E-22	188	67	35
NP_249783	488	167	9.00E-41	242	103	42	AAP78393	514	68.9	5.00E-11	115	40	34
NP_249783	488	113	2.00E-24	335	100	29	NP_420273	273	104	1.00E-21	297	76	25
ZP_00138680488	167	9.00E-41	242	103	42	NP_907654	518	103	2.00E-21	192	63	32	
ZP_00138680488	113	2.00E-24	335	100	29	NP_907654	518	94.7	8.00E-19	333	85	25	
ZP_00130923288	167	9.00E-41	284	103	36	NP_404388	404	102	3.00E-21	223	67	30	
NP_798637	376	167	1.00E-40	377	125	33	NP_670739	401	102	3.00E-21	223	67	30
NP_798638	378	167	1.00E-40	377	126	33	NP_670739	401	57.4	1.00E-07	84	28	33
AAP08637	249	165	5.00E-40	271	98	36	NP_282484	572	102	5.00E-21	181	66	35
ZP_00317212530	184	6.00E-40	222	101	45	NP_282484	572	72.8	3.00E-12	132	45	34	
ZP_00317212530	99.4	3.00E-20	182	66	36	NP_214372	518	100	1.00E-20	206	62	29	
NP_746492	687	164	8.00E-40	252	103	40	NP_214372	518	62	6.00E-09	129	39	30
NP_746492	687	96.3	3.00E-19	129	53	41	ZP_00313281302	99.4	3.00E-20	274	69	25	
ZP_00100062385	163	1.00E-39	386	128	33	ZP_00100043955	99.4	3.00E-20	270	82	30		
ZP_00317210587	162	4.00E-39	254	105	41	ZP_00100043955	63.5	2.00E-09	154	47	30		
ZP_00317210587	95.9	3.00E-19	129	55	42	NP_420274	273	99	4.00E-20	292	75	25	
NP_712200	283	162	4.00E-39	301	100	33	NP_419609	273	99	4.00E-20	292	72	24
YP_001838	283	161	5.00E-39	301	99	32	NP_419611	273	98.6	5.00E-20	292	72	24
NP_770335	274	160	1.00E-38	295	95	32	YP_071817	400	98.2	7.00E-20	207	61	29
ZP_00218998614	156	2.00E-37	242	97	40	YP_071817	400	60.1	2.00E-06	115	37	32	
ZP_00218998614	90.1	2.00E-17	86	46	53	NP_865696	718	98.2	7.00E-20	284	88	30	
ZP_00270146298	154	1.00E-36	305	107	35	NP_865696	718	81.3	9.00E-15	143	57	39	
NP_935285	396	153	1.00E-36	376	118	31	NP_622171	296	98.2	7.00E-20	290	83	28
NP_945993	274	152	3.00E-36	295	92	31	AAP78250	508	97.8	9.00E-20	170	60	35
NP_759224	377	151	5.00E-36	376	120	31	AAP78250	508	76.3	3.00E-13	297	74	24
NP_933767	377	151	7.00E-36	376	119	31	NP_419610	273	97.4	1.00E-18	292	70	23
YP_128298	426	151	7.00E-36	251	106	42	NP_866107	739	97.4	1.00E-19	257	76	29
YP_128298	89.4	3.00E-17	149	56	36	NP_866107	739	79.3	3.00E-14	250	70	28	
ZP_00329901380	149	2.00E-36	382	125	32	NP_223266	510	96.3	3.00E-19	170	57	33	
NP_935286	377	149	2.00E-35	376	112	29	NP_223265	510	82.4	4.00E-15	311	78	25
NP_933764	385	149	3.00E-35	178	80	44	NP_404369	399	95.9	3.00E-19	295	76	25
NP_933764	385	102	5.00E-21	301	92	30	ZP_00299642294	95.5	5.00E-19	295	75	25	
ZP_00273972629	146	2.00E-34	199	85	42	NP_867573	685	92.8	3.00E-18	189	62	36	
ZP_00273972629	104	8.00E-22	92	57	61	NP_867573	685	85.5	5.00E-16	126	55	43	
YP_129126	393	145	4.00E-34	213	82	38	NP_404370	401	92	5.00E-18	309	79	25
YP_129126	90.5	1.00E-17	203	63	31	ZP_00329910305	91.7	7.00E-18	285	73	25		
NP_978099	266	145	4.00E-34	292	96	32	ZP_00196929279	91.7	7.00E-18	288	79	27	
ZP_00287990272	142	4.00E-33	290	85	30	NP_353572	306	91.3	9.00E-18	313	77	24	
YP_035878	266	142	4.00E-33	292	97	33	NP_693427	293	91.3	9.00E-18	293	75	25
NP_759225	374	141	6.00E-33	167	76	45	ZP_0008298321	90.9	1.00E-17	302	85	28	
NP_759225	374	102	5.00E-21	301	92	30	NP_227947	258	89.7	3.00E-17	243	63	25
NP_782289	268	141	6.00E-33	266	79	29	NP_954084	298	89.7	3.00E-17	290	75	25
ZP_00004597493	140	9.00E-33	166	74	44	ZP_00289010110	89.7	3.00E-17	87	47	54		
ZP_00004597493	97.1	2.00E-19	176	63	35	NP_700301	282	88.6	6.00E-17	298	76	25	
YP_083129	266	140	9.00E-33	292	95	32	NP_391420	298	86.7	2.00E-16	288	80	27
AAP08634	266	140	1.00E-32	291	95	32	NP_541127	282	86.7	2.00E-16	298	75	25
NP_228567	387	140	2.00E-32	392	119	30	ZP_00288130268	85.5	5.00E-16	295	76	25	
ZP_00235456266	138	5.00E-32	292	96	32	NP_773505	313	85.1	6.00E-16	311	73	23	
YP_083130	460	137	8.00E-32	289	96	33	NP_104150	328	84.7	8.00E-16	329	75	22
YP_083130	460	80.9	1.00E-14	111	46	41	ZP_00339045281	84	1.00E-15	296	75	25	
NP_342261	269	137	1.00E-31	292	89	30	YP_080866	303	83.2	2.00E-15	281	70	24
NP_242343	464	130	2.00E-29	135	70	51	NP_773505	314	80.9	1.00E-14	317	73	23
NP_242343	464	95.9	3.00E-19	169	57	33	NP_353570	313	79.7	3.00E-14	313	75	23
NP_978100	465	126	2.00E-28	221	77	34	ZP_00339044282	79.3	3.00E-14	298	72	24	
NP_978100	465	75.1	6.00E-13	84	41	NP_907272	253	74.3	1.00E-12	278	65	23	
AAO08750	375	121	6.00E-27	374	108	28	NP_840399	299	77.8	1.00E-13	290	71	24
NP_933768	375	121	6.00E-27	374	108	28	ZP_00339046273	77.4	1.00E-13	280	73	26	
NP_391395	160	116	2.00E-25	179	68	37	ZP_00338034282	77.4	1E-13	294	72	24	
NP_797170	374	109	2.00E-23	148	60	40	ZP_00193517283	76.5	2.00E-13	290	65	22	
NP_797170	374	74.3	1.00E-12	217	61	28	ZP_00337030282	76.3	3.00E-13	299	69	23	
NP_908288	513	109	2.00E-23	294	89	30	NP_907272	253	74.3	1.00E-12	278	65	23
NP_908288	513	73.2	2.00E-12	122	44	36	ZP_00270458289	72.8	3.00E-12	302	69	22	
NP_208915	514	108	7.00E-23	187	67	35	NP_00289016275	72	5.00E-12	275	64	23	
NP_208915	514	77.4	1.00E-13	121	47	38	NP_353571	337	71.6	7.00E-12	324	75	23
NP_222828	514	107	1.00E-22	187	67	35	NP_531244	320	71.6	7.00E-12	324	75	23
NP_222828	514	77.4	1.00E-13	121	47	38	ZP_00183364320	71.6	7E-12	321	74	23	

Fig. 25C-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
NP_281892	248	68.6	6.00E-11	288	66	22	ZP_00317213517	53.1	3.00E-06	180	31	31	
NP_782319	316	68.2	8.00E-11	284	68	23	ZP_00317213517	40	2.30E-02	91	24	26	
ZP_00337274271	67.4	1.00E-10	284	54	22	ZP_00273669412	52.8	3.00E-06	311	67	21		
ZP_00335243300	67	2.00E-10	302	74	24	ZP_00273669412	35.8	4.30E-01	146	35	23		
ZP_00149794326	67	2.00E-10	322	67	20	ZP_00056435521	52.8	3.00E-06	185	40	21		
NP_972952	415	66.6	2.00E-10	285	65	22	NP_384777	401	52.8	3.00E-06	189	48	25
NP_227898	304	65.1	7.00E-10	256	62	24	NP_384777	401	40	2.30E-02	87	24	27
YP_013346	291	64.3	1.00E-09	288	61	21	NP_415601	317	52	6.00E-06	305	70	22
ZP_00049388242	63.9	1.00E-09	188	52	27	NP_421770	424	52	6.00E-06	184	35	21	
NP_464233	291	63.9	1.00E-09	238	61	21	NP_287217	317	51.6	3.00E-06	299	66	22
ZP_00229464291	63.5	2.00E-09	238	61	21	NP_967522	361	51.6	8.00E-06	178	44	24	
AAC65633	416	63.2	3.00E-09	268	60	22	YP_068426	757	51.6	8.00E-06	130	40	30
NP_384778	321	62.4	4.00E-09	321	68	21	YP_111446	2634	51.2	1.00E-05	287	60	20
NP_642302	401	62.4	4.00E-09	134	40	29	YP_111446	2634	50.8	1.00E-05	281	59	20
NP_642302	401	40.8	1.30E-02	118	31	26	YP_111446	2634	47	2.00E-04	296	63	21
NP_470057	291	62	6.00E-09	288	60	20	YP_111446	2634	46.2	3.00E-04	291	60	20
ZP_00237150287	60.8	1.00E-08	243	52	25	YP_111446	2634	45.1	7.00E-04	292	58	19	
NP_348828	425	60.1	2.00E-08	289	65	22	YP_111446	2634	43.5	2.00E-03	284	58	20
NP_348828	425	42.4	5.00E-03	310	53	22	YP_111446	2634	43.5	2.00E-03	283	58	20
YP_126585	411	60.1	2.00E-08	147	39	26	YP_111446	2634	43.5	2.00E-03	289	59	20
YP_095257	411	60.1	2.00E-08	147	39	26	YP_111446	2634	43.1	3.00E-03	292	59	20
NP_104151	356	59.7	3.00E-08	357	81	22	YP_111446	2634	42	6.00E-03	281	63	22
YP_123658	411	59.3	4.00E-08	147	38	25	YP_111446	2634	40	2.30E-02	295	59	20
YP_123658	411	32	6.20E+00	83	25	30	YP_111446	2634	40	2.30E-02	298	59	19
AAQ60546	309	58.9	5.00E-08	299	62	20	YP_111446	2634	39.3	3.90E-02	302	59	19
ZP_00127287530	58.9	5.00E-08	135	40	29	YP_111446	2634	38.9	5.10E-02	283	53	18	
NP_772484	757	58.5	6.00E-08	273	72	26	YP_111446	2634	37.4	1.50E-01	278	48	17
NP_772484	757	34.7	9.60E-01	109	29	26	YP_111446	2634	35.8	4.30E-01	235	50	21
NP_537307	401	58.2	8.00E-08	134	38	28	YP_105520	459	51.2	1.00E-05	246	57	22
NP_537307	401	35.8	4.30E-01	119	31	26	YP_105520	459	41.6	8.00E-03	298	60	20
YP_083110	287	57.8	1.00E-07	242	57	23	YP_027819	207	51.2	1.00E-05	115	30	26
NP_978079	287	57.8	1.00E-07	241	60	24	NP_866849	651	51.2	1.00E-05	129	34	26
NP_772486	757	57.4	1.00E-07	265	72	28	NP_947642	935	50.8	1.00E-05	284	70	24
NP_772486	757	34.3	1.20E+00	109	29	26	NP_947642	935	34.3	1.20E+00	93	22	23
NP_212316	424	57.4	1.00E-07	132	38	28	NP_949286	888	50.8	1.00E-05	284	68	24
NP_212316	424	35	7.30E-01	77	18	23	NP_949286	888	38.9	5.10E-02	305	63	20
NP_384775	394	57	2.00E-07	138	38	27	YP_009743	523	50.4	2.00E-05	123	32	26
NP_384775	394	60.4	2.00E-05	87	29	33	YP_009743	523	33.9	1.60E+00	171	36	21
YP_072631	424	56.6	2.00E-07	157	46	29	ZP_00129084282	50.4	2.00E-05	285	68	23	
YP_072631	424	35	7.30E-01	77	18	23	NP_282040	750	50.1	2.00E-05	129	36	27
AAP08615	287	56.6	2.00E-07	243	57	23	NP_282040	750	41.6	8.00E-03	228	48	26
ZP_00128974504	56.6	2.00E-07	113	32	28	ZP_00242042398	49.7	3.00E-05	215	53	24		
ZP_00128974504	43.5	2.00E-03	238	50	21	ZP_00242042398	39.7	3.00E-02	140	35	25		
NP_521913	316	56.2	3.00E-07	305	65	21	ZP_00263356523	48.5	6.00E-05	162	43	26	
NP_746494	521	56.2	3.00E-07	130	41	31	ZP_00263356523	35	7.30E-01	106	23	21	
NP_746494	521	41.6	8.00E-03	149	37	24	NP_460155	317	48.5	6.00E-05	140	38	27
ZP_00173446304	56.2	3.00E-07	301	68	22	YP_128296	304	48.5	6.00E-05	293	53	18	
ZP_00278994407	55.5	5.00E-07	250	55	22	YP_040849	10746	48.5	6.00E-05	303	63	20	
NP_718794	403	55.1	7.00E-07	136	36	26	YP_040849	10746	42.4	6.00E-03	340	71	20
NP_718794	403	34.3	1.20E+00	89	26	29	YP_040849	10746	36.6	2.60E-01	297	58	19
NP_791768	530	55.1	7.00E-07	130	36	27	YP_040849	10746	36.6	2.50E-01	332	69	20
NP_772485	757	55.1	7.00E-07	250	63	25	YP_040849	10746	35.8	4.30E-01	301	63	20
NP_772485	757	33.9	1.60E+00	109	29	26	YP_040849	10746	33.5	2.10E+00	279	56	19
NP_880130	510	54.7	9.00E-07	136	43	31	YP_040849	10746	33.1	2.80E+00	317	68	21
YP_003353	422	54.7	9.00E-07	110	34	30	YP_040849	10746	31.6	8.10E+00	297	57	19
NP_892941	337	54.3	1.00E-06	310	73	23	ZP_00167907404	48.5	6.00E-05	121	37	30	
NP_883789	510	54.3	1.00E-06	136	42	30	ZP_00167907404	36.4	5.60E-01	99	23	23	
NP_405374	326	54.3	1.00E-06	310	73	23	NP_384775	394	47.8	1.00E-04	137	34	24
ZP_00212998398	54.3	1.00E-06	111	29	26	NP_384776	394	45.8	4.00E-04	245	56	22	
NP_799784	299	53.9	2.00E-06	258	55	21	NP_562800	451	47.4	1.00E-04	220	53	24
NP_772483	763	53.9	2.00E-06	186	47	25	NP_455577	317	47.4	1.00E-04	140	38	27
NP_772483	763	36.2	3.30E-01	109	29	26	NP_646141	9904	47	2.00E-04	306	64	20
NP_244487	395	53.9	2.00E-06	126	33	26	NP_646141	9904	43.1	3.00E-03	153	36	23
NP_353593	436	53.5	2.00E-06	256	59	23	NP_646141	9904	39.7	3.00E-02	332	71	21
NP_353593	436	46.2	3.00E-04	140	34	24	NP_646141	9904	39.3	3.90E-02	283	66	23
NP_753263	317	53.6	2.00E-06	299	67	22	NP_646141	9904	38.1	8.60E-02	279	56	20
NP_531268	430	53.5	2.00E-06	256	59	23	NP_646141	9904	35.4	5.60E-01	317	66	20
NP_531268	430	46.2	3.00E-04	140	34	24	NP_646141	9904	33.9	1.60E+00	297	57	19

Fig. 25C-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
YP_012160	282	47	2.00E-04	223	53	23	YP_105401	1535	42.4	5.00E-03	300	63	21
NP_249778	439	46.6	2.00E-04	134	34	25	YP_105401	1535	36.2	3.30E-01	307	84	20
NP_249778	439	32	6.20E+00	257	50	19	YP_105401	1535	35.8	4.30E-01	217	46	21
ZP_00138575439	46.6	2.00E-04	134	34	25	YP_065260	567	42.4	5.00E-03	263	54	20	
ZP_00314296822	46.2	3.00E-04	257	54	21	NP_905736	846	42.4	5.00E-03	168	37	22	
NP_223759	267	46.2	3.00E-04	148	40	26	ZP_001432362005	42.4	5.00E-03	255	59	23	
YP_108306	1806	46.2	3.00E-04	296	66	22	NP_523179	1309	42	6.00E-03	264	60	22
YP_108306	1606	38.1	8.60E-02	281	57	20	NP_523179	1309	36.2	3.30E-01	250	56	21
YP_108306	1806	38.1	8.60E-02	228	54	23	NP_523179	1309	35.8	4.30E-01	247	57	23
YP_108306	1606	38.1	8.60E-02	309	60	19	AAF95046	672	42	6.00E-03	320	64	20
YP_108306	1606	37.4	1.50E-01	256	58	22	NP_468975	661	42	6.00E-03	268	63	23
ZP_00290372765	45.8	4.00E-04	280	56	20	NP_784110	1377	42	6.00E-03	285	64	22	
NP_267773	1072	45.4	5.00E-04	287	53	18	YP_129124	396	41.6	8.00E-03	119	25	21
NP_267773	1072	33.5	2.10E+00	91	21	23	ZP_00069080401	41.6	8.00E-03	122	36	29	
NP_228520	407	45.4	5.00E-04	290	63	21	ZP_00089080401	38.5	6.60E-02	139	36	25	
ZP_003230792334	45.4	5.00E-04	274	50	18	YP_465811	1787	41.2	1.00E-02	266	58	21	
ZP_003230792334	39.7	3.00E-02	276	49	17	ZP_002143701439	41.2	1.00E-02	297	72	24		
ZP_003230792334	39.3	3.90E-02	293	54	18	ZP_002143701439	32.3	4.70E+00	272	56	20		
ZP_003230792334	38.5	6.60E-02	294	53	18	AAP07456	953	40.8	1.30E-02	263	49	18	
ZP_003230792334	37	1.90E-01	280	52	18	ZP_00063096901	40.8	1.30E-02	241	60	24		
ZP_003230792334	34.7	9.60E-01	261	50	19	ZP_00296669699	40.4	1.70E-02	292	54	18		
YP_111449	1530	45.1	7.00E-04	295	64	21	ZP_00296669699	32	6.20E+00	133	26	19	
YP_111449	1530	44.7	1.00E-03	292	64	21	NP_251152	5627	40.4	1.70E-02	329	67	20
YP_111449	1630	44.3	1.00E-03	278	59	21	NP_251152	5627	34.3	1.20E+00	235	50	21
YP_111449	1530	42.7	4.00E-03	320	69	21	NP_251152	5627	32.7	3.60E+00	312	60	19
YP_111449	1530	41.6	8.00E-03	288	59	20	ZP_002886079749	40	2.30E-02	236	51	21	
YP_111449	1530	41.2	1.00E-02	284	60	21	YP_142319	789	40	2.30E-02	302	64	21
YP_111449	1530	40	2.30E-02	250	51	20	YP_140402	710	40	2.30E-02	302	64	21
YP_111449	1530	37.7	1.10E-01	250	50	20	NP_373178	2271	40	2.30E-02	287	52	18
YP_111449	1530	37	1.90E-01	263	52	19	NP_373178	2271	34.3	1.20E+00	216	42	19
YP_111449	1530	34.7	9.60E-01	257	55	21	NP_373178	2271	32	6.20E+00	303	62	20
NP_949252	886	44.7	1.00E-03	194	49	25	AAO08483	623	40	2.30E-02	114	31	27
NP_949252	886	33.9	1.60E+00	102	25	24	YP_114336	437	40	2.30E-02	83	27	32
NP_929192	321	44.7	1.00E-03	318	74	23	YP_034700	953	40	2.30E-02	301	58	19
NP_647392	2275	44.7	1.00E-03	287	53	18	NP_936493	623	40	2.30E-02	114	31	27
NP_647392	2275	34.3	1.20E+00	216	42	19	NP_764683	8439	40	2.30E-02	130	27	20
NP_647392	2275	34.3	1.20E+00	299	60	20	NP_764683	8439	38.5	6.60E-02	169	43	25
NP_371958	6713	44.7	1.00E-03	303	61	20	NP_764683	8439	36.2	3.30E-01	265	55	20
NP_371958	6713	42	6.00E-03	153	36	23	NP_764683	8439	36.2	3.30E-01	288	61	21
NP_371958	6713	41.2	1.00E-02	332	72	21	NP_764683	8439	36.2	3.30E-01	265	48	18
NP_371958	6713	35.4	5.60E-01	276	53	19	NP_764683	8439	35.4	5.60E-01	316	61	19
NP_371958	6713	33.1	2.80E+00	297	57	19	NP_764683	8439	34.7	9.60E-01	302	60	19
NP_374548	6713	44.7	1.00E-03	303	61	20	NP_764683	8439	33.9	1.60E+00	314	65	20
NP_374548	6713	42	6.00E-03	153	36	23	NP_764683	8439	33.9	1.60E+00	146	40	27
NP_374548	6713	41.2	1.00E-02	332	72	21	NP_764683	8439	33.1	2.80E+00	181	46	25
NP_374548	6713	36.4	5.60E-01	276	53	19	NP_764683	8439	33.1	2.80E+00	151	38	25
NP_374548	6713	33.1	2.80E+00	297	57	19	NP_764683	8439	32.3	4.70E+00	329	68	20
YP_106908	410	44.7	1.00E-03	124	28	22	NP_764683	8439	31.6	8.10E+00	292	53	18
YP_044654	2275	44.7	1.00E-03	287	53	18	NP_00123591638	39.7	3.00E-02	352	74	21	
YP_044654	2275	34.3	1.20E+00	216	42	19	ZP_000642471003	39.7	3.00E-02	307	71	23	
YP_110805	1653	44.3	1.00E-03	269	58	21	ZP_00319871233	39.3	3.90E-02	139	37	26	
YP_110805	1653	40	2.30E-02	254	56	22	NP_908061	118	39.3	3.90E-02	103	29	28
YP_110805	1653	37	1.90E-01	287	58	20	ZP_002335911787	39.3	3.90E-02	261	60	22	
YP_110805	1653	36.2	3.30E-01	274	56	20	NP_798012	3240	38.9	5.10E-02	269	58	21
YP_110805	1653	34.7	9.60E-01	265	62	19	YP_015633	662	38.9	5.10E-02	167	43	25
ZP_001527551140	44.3	1.00E-03	295	62	21	ZP_002814314726	38.9	5.10E-02	310	62	20		
AAAF94608	4558	43.9	2.00E-03	228	52	22	ZP_002814314726	38.1	8.60E-02	308	61	19	
ZP_00326048609	43.1	3.00E-03	273	63	23	ZP_002814314726	37.4	1.50E-01	272	54	19		
ZP_00326048609	32	6.20E+00	99	20	20	ZP_002814314726	36.6	2.50E-01	275	56	20		
ZP_00051799172	43.1	3.00E-03	136	31	22	ZP_002814314726	36.6	2.50E-01	254	53	20		
NP_207286	295	43.1	3.00E-03	150	42	28	ZP_002814314726	35.4	5.60E-01	279	55	19	
NP_936893	989	43.1	3.00E-03	289	63	21	ZP_002814314726	35	7.30E-01	319	59	18	
AAQ59385	302	42.7	4.00E-03	282	55	19	ZP_002814314726	34.3	1.20E+00	324	59	18	
NP_267008	1063	42.7	4.00E-03	291	56	18	ZP_00211493472	38.9	6.10E-02	274	60	21	
NP_267008	1063	37	1.90E-01	231	42	18	ZP_002047941979	38.9	5.10E-02	235	47	20	
NP_267008	1063	36.6	2.50E-01	281	63	22	ZP_002047941979	34.7	9.60E-01	236	50	21	
YP_042074	1351	42.7	4.00E-03	304	59	19	ZP_002047941979	33.1	2.8	312	61	19	
NP_939992	1254	42.4	5.00E-03	189	38	22	ZP_00206729459	38.5	6.60E-02	278	63	22	

Fig. 25C-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
NP_928380	567	38.5	6.60E-02	257	58	22	ZP_00160517717	31.6	8.10E+00	248	46	18	
YP_115646	760	38.5	6.60E-02	144	32	22	ZP_001380311417	36.2	3.30E-01	201	49	24	
YP_087940	5399	38.5	6.60E-02	312	72	23	NP_561526	1109	35.8	4.30E-01	287	53	18
YP_087940	5399	37.4	1.50E-01	256	45	17	NP_561526	1109	32	6.20E+00	181	40	22
YP_087940	5399	37.4	1.50E-01	256	45	17	NP_349224	664	35.8	4.30E-01	272	59	21
YP_087940	5399	36.2	3.30E-01	273	58	21	AAP56614	499	35.8	4.30E-01	183	40	21
YP_087940	5399	36.2	3.30E-01	273	58	21	YP_133618	665	35.8	4.30E-01	111	33	29
YP_087940	5399	35.8	4.30E-01	274	51	22	YP_071943	3378	35.8	4.30E-01	289	61	21
YP_087940	5399	35	7.30E-01	294	58	19	ZP_00270453727	35.4	5.60E-01	239	49	20	
YP_087940	5399	33.9	1.60E+00	289	57	19	AAQ59977	458	35.4	5.60E-01	266	61	22
YP_087940	5399	33.5	2.10E+00	289	57	19	NP_773497	491	35.4	5.60E-01	121	26	23
NP_781518	516	38.5	6.60E-02	302	64	21	NP_267038	799	35.4	5.60E-01	301	71	23
ZP_00213493557	38.5	6.60E-02	223	51	22	NP_478503	1487	35.4	5.60E-01	249	49	19	
ZP_00224552561	38.5	6.60E-02	223	53	23	YP_086339	664	35.4	5.60E-01	107	23	21	
ZP_00355727801	38.5	6.60E-02	317	71	22	NP_207948	1230	35.4	5.60E-01	292	62	21	
ZP_001404443443	38.5	6.60E-02	286	62	21	YP_039058	564	35.4	5.60E-01	107	23	21	
NP_419715	307	38.1	8.60E-02	73	21	NP_864533	7715	35.4	5.60E-01	240	52	21	
NP_207906	228	38.1	8.60E-02	110	32	29	NP_765804	2310	35.4	5.60E-01	290	52	17
ZP_003235731130	38.1	8.60E-02	292	65	22	NP_765787	676	35.4	5.60E-01	117	33	28	
ZP_003235731130	32.3	4.70E+00	281	57	20	ZP_00323298769	35.4	5.60E-01	148	35	23		
ZP_00317192676	37.7	1.10E-01	169	35	20	ZP_002889038997	36.4	5.60E-01	294	56	19		
NP_771092	696	37.7	1.10E-01	134	39	29	ZP_00338261539	35	7.30E-01	62	20	32	
NP_214145	422	37.7	1.10E-01	186	40	21	NP_718836	667	35	7.30E-01	98	20	20
NP_766958	681	37.7	1.10E-01	176	45	25	YP_125297	361	35	7.30E-01	368	69	19
ZP_003125671479	37.7	1.10E-01	273	58	21	YP_132326	529	35	7.30E-01	332	69	20	
ZP_00240971654	37.7	1.10E-01	107	24	22	NP_812240	868	35	7.30E-01	150	44	29	
ZP_001952636375	37.7	1.10E-01	183	41	22	NP_964984	4734	36	7.30E-01	316	66	20	
NP_949268	623	37.4	1.50E-01	277	53	19	NP_964984	4734	33.1	2.80E+00	153	40	26
ZP_00315446537	37.4	1.50E-01	263	54	20	ZP_00144637646	35	7.30E-01	243	44	18		
ZP_0028945115245	37.4	1.50E-01	272	65	23	ZP_00222048485	35	7.30E-01	88	23	26		
ZP_00273578527	37.4	1.50E-01	90	25	27	ZP_00220113501	35	7.30E-01	88	23	26		
ZP_00273578527	33.9	1.60E+00	83	22	NP_00137782529	35	7.30E-01	245	43	17			
NP_800021	623	37.4	1.50E-01	117	31	NP_00121836231	35	7.30E-01	165	39	23		
NP_772493	627	37.4	1.50E-01	229	49	21	NP_946001	587	34.7	9.60E-01	121	36	29
AAP78292	855	37.4	1.50E-01	138	32	23	ZP_00319892754	34.7	9.60E-01	284	62	21	
ZP_00281310507	37.4	1.50E-01	119	31	26	ZP_00319892754	33.1	2.80E+00	261	51	19		
ZP_00289133779	37	1.90E-01	198	38	19	ZP_00311897602	34.7	9.60E-01	269	55	20		
ZP_00289133779	36.2	3.30E-01	89	26	29	NP_00336529824	34.7	9.60E-01	149	35	23		
NP_471790	927	37	1.90E-01	281	59	20	NP_802026	593	34.7	9.60E-01	118	33	27
NP_471790	927	31.6	8.10E+00	210	46	21	NP_924800	1241	34.7	9.60E-01	73	24	32
NP_522834	3322	37	1.90E-01	228	52	22	NP_463823	500	34.7	9.60E-01	233	47	20
NP_404364	307	37	1.90E-01	253	52	20	NP_645681	946	34.7	9.60E-01	219	44	20
NP_336479	342	37	1.90E-01	87	25	28	NP_792425	633	34.7	9.60E-01	283	50	17
NP_248731	3535	37	1.90E-01	286	62	21	NP_561978	344	34.7	9.60E-01	232	49	21
YP_066441	693	37	1.90E-01	120	30	25	NP_350172	783	34.7	9.60E-01	129	29	22
NP_757754	161	37	1.90E-01	77	23	NP_349191	722	34.7	9.60E-01	143	36	25	
NP_670743	308	37	1.90E-01	253	52	20	NP_349191	722	33.1	2.80E+00	292	57	19
NP_603291	1724	37	1.90E-01	131	34	25	AAO10484	626	34.7	9.60E-01	299	52	17
ZP_00290809010	37	1.90E-01	260	53	20	NP_642322	396	34.7	9.60E-01	123	22	17	
ZP_00219646252	37	1.90E-01	74	21	28	YP_133410	577	34.7	9.60E-01	317	51	19	
ZP_00053136721	37	1.90E-01	298	58	19	NP_265927	594	34.7	9.60E-01	118	33	27	
NP_799785	346	36.6	2.50E-01	116	29	NP_042878	928	34.7	9.60E-01	219	44	20	
NP_739300	290	36.6	2.50E-01	122	30	YP_012922	500	34.7	9.60E-01	233	47	20	
NP_996362	1119	36.6	2.50E-01	143	35	24	NP_051325	222	34.7	9.60E-01	114	30	26
NP_359655	1902	36.6	2.50E-01	282	60	21	NP_782459	569	34.7	9.60E-01	305	61	20
NP_253231	1417	36.6	2.50E-01	201	49	24	NP_607532	594	34.7	9.60E-01	118	33	27
NP_932208	603	36.6	2.50E-01	53	17	NP_00233975541	34.7	9.60E-01	233	47	20		
ZP_0021898486	36.6	2.50E-01	259	56	21	ZP_00212291306	34.7	9.60E-01	130	38	29		
ZP_00149774452	36.6	2.50E-01	264	56	21	NP_469464	1788	34.3	1.20E+00	290	65	22	
ZP_00319246821	36.2	3.30E-01	65	19	NP_792292	629	34.3	1.20E+00	128	31	24		
ZP_00284479601	36.2	3.30E-01	262	52	19	NP_695350	458	34.3	1.20E+00	271	54	18	
ZP_0005278425	36.2	3.30E-01	193	50	25	NP_348729	570	34.3	1.20E+00	162	36	22	
NP_280332	643	36.2	3.30E-01	233	50	21	NP_252997	632	34.3	1.20E+00	245	43	17
NP_522101	3552	36.2	3.30E-01	209	45	21	YP_066477	393	34.3	1.20E+00	148	33	22
NP_964415	1096	36.2	3.30E-01	265	56	21	YP_064761	880	34.3	1.20E+00	126	30	23
ZP_00313063880	36.2	3.30E-01	181	39	21	YP_064723	674	34.3	1.20E+00	248	48	19	
ZP_00182337425	36.2	3.30E-01	245	41	16	NP_935137	626	34.3	1.20E+00	299	52	17	
ZP_00160617717	36.2	3.30E-01	289	62	21	NP_866060	3056	34.3	1.20E+00	120	34	28	

Fig. 25C-Continued

Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S	Hit_ID	Length	Bitscore	Expected	Length	#Ident	%S
NP_866060_3056	33.1	2.80E+00	116	34	29		ZP_00147410749	33.1	2.80E+00	131	34	25	
NP_616105_1074	34.3	1.20E+00	257	49	19		ZP_00137783633	33.1	2.80E+00	246	42	17	
ZP_003233981656	34.3	1.20E+00	187	41	21		ZP_00123950634	33.1	2.80E+00	96	24	25	
ZP_00308827366	34.3	1.20E+00	138	35	25		ZP_00128148551	33.1	2.80E+00	256	46	17	
ZP_00137780616	34.3	1.20E+00	245	43	17		ZP_00326431563	33.1	2.80E+00	274	69	25	
ZP_00124017629	34.3	1.20E+00	254	45	18		ZP_00326431563	32.7	3.60E+00	307	70	22	
ZP_00110825475	34.3	1.20E+00	140	35	25		NP_252999_629	32.7	3.60E+00	127	32	25	
ZP_00046942967	33.9	1.60E+00	134	35	26		ZP_000531851136	32.7	3.60E+00	223	51	22	
YP_080405_660	33.9	1.60E+00	87	22	25		NP_993534_2578	32.7	3.60E+00	280	61	21	
NP_470074_599	33.9	1.60E+00	288	69	20		YP_081346_561	32.7	3.60E+00	99	20	20	
NP_927898_4582	33.9	1.60E+00	257	61	23		NP_471252_1186	32.7	3.60E+00	315	63	20	
NP_815456_533	33.9	1.60E+00	125	35	28		NP_469482_290	32.7	3.60E+00	82	21	25	
NP_562046_633	33.9	1.60E+00	259	51	19		NP_464250_601	32.7	3.60E+00	193	39	20	
NP_267826_901	33.9	1.60E+00	281	56	19		NP_929013_190	32.7	3.60E+00	133	34	25	
YP_135620_471	33.9	1.60E+00	121	30	24		NP_793014_6274	32.7	3.60E+00	246	51	20	
YP_126425_657	33.9	1.80E+00	121	29	23		NP_790315_541	32.7	3.60E+00	93	28	30	
YP_131301_638	33.9	1.60E+00	73	16	21		NP_769573_564	32.7	3.60E+00	157	36	22	
NP_987906_720	33.9	1.60E+00	95	27	28		NP_563507_721	32.7	3.60E+00	112	26	22	
YP_030040_660	33.9	1.60E+00	105	25	23		NP_406024_2535	32.7	3.60E+00	290	61	21	
NP_981473_564	33.9	1.60E+00	109	24	22		NP_939703_888	32.7	3.60E+00	123	31	25	
NP_784971_983	33.9	1.60E+00	299	58	19		AAP56776_742	32.7	3.60E+00	186	40	21	
NP_783077_417	33.9	1.60E+00	103	24	23		NP_882071_1175	32.7	3.60E+00	65	18	27	
NP_965011_912	33.9	1.60E+00	214	49	22		NP_419323_622	32.7	3.60E+00	235	47	20	
NP_964462_1000	33.9	1.60E+00	125	28	22		YP_134634_596	32.7	3.60E+00	63	14	22	
ZP_002891151200	33.5	2.10E+00	61	17	27		YP_108428_634	32.7	3.60E+00	266	52	19	
ZP_00056271661	33.5	2.10E+00	278	54	19		NP_109825_228	32.7	3.60E+00	154	29	18	
NP_287395_973	33.5	2.10E+00	185	42	22		YP_013363_601	32.7	3.60E+00	193	39	20	
NP_794375_539	33.5	2.10E+00	128	23	17		NP_669014_2579	32.7	3.60E+00	290	61	21	
YP_137657_727	33.5	2.10E+00	86	25	29		ZP_00284287481	32.7	3.60E+00	77	21	27	
YP_115527_534	33.5	2.10E+00	80	24	30		ZP_D0280424428	32.7	3.60E+00	186	50	25	
NP_309677_971	33.5	2.10E+00	185	42	22		NP_326456_750	32.3	4.70E+00	121	27	22	
NP_936395_542	33.5	2.10E+00	300	62	20		NP_391003_662	32.3	4.70E+00	322	60	18	
NP_637327_396	33.5	2.10E+00	123	21	17		NP_852542_1658	32.3	4.70E+00	149	33	22	
NP_616912_1052	33.5	2.10E+00	132	36	27		NP_852542_1658	32.3	4.70E+00	139	27	19	
NP_984155_982	33.5	2.10E+00	203	47	23		NP_646487_424	32.3	4.70E+00	147	36	24	
NP_00296785641	33.5	2.10E+00	275	60	21		NP_718941_169	32.3	4.70E+00	160	36	22	
ZP_00217082977	33.5	2.10E+00	109	32	29		NP_716901_706	32.3	4.70E+00	273	59	21	
ZP_00106667978	33.5	2.10E+00	112	29	25		NP_801002_542	32.3	4.70E+00	304	58	19	
ZP_00290912684	33.1	2.80E+00	130	24	18		AAQ60540_2373	32.3	4.70E+00	255	55	21	
ZP_00290912684	32.3	4.70E+00	188	35	18		AAQ58389_690	32.3	4.70E+00	110	25	22	
ZP_003382277837	33.1	2.80E+00	109	26	23		NP_770343_582	32.3	4.70E+00	270	50	18	
NP_798538_678	33.1	2.80E+00	127	27	21		AAF66820_652	32.3	4.70E+00	120	31	25	
NP_790701_498	33.1	2.80E+00	162	41	25		AAF94794_596	32.3	4.70E+00	80	19	23	
NP_768659_816	33.1	2.80E+00	62	19	30		NP_404241_3295	32.3	4.70E+00	289	61	21	
NP_415890_1122	33.1	2.80E+00	147	37	25		YP_137110_536	32.3	4.70E+00	199	47	23	
AAQ08482_622	33.1	2.80E+00	120	30	25		YP_032604_1872	32.3	4.70E+00	118	31	26	
AAQ10918_675	33.1	2.80E+00	125	29	23		NP_782523_471	32.3	4.70E+00	107	20	18	
NP_889112_532	33.1	2.80E+00	250	48	19		NP_785908_300	32.3	4.70E+00	132	29	21	
YP_129467_640	33.1	2.80E+00	137	31	22		NP_785785_952	32.3	4.70E+00	195	37	18	
YP_133515_467	33.1	2.80E+00	127	26	20		NP_506640_628	32.3	4.70E+00	271	60	22	
YP_132040_542	33.1	2.80E+00	107	27	25		ZP_002394591038	32.3	4.70E+00	243	52	21	
YP_076807_558	33.1	2.80E+00	118	30	25		ZP_00236267289	32.3	4.70E+00	156	31	19	
NP_987807_729	33.1	2.80E+00	95	26	27		ZP_002326811086	32.3	4.70E+00	277	57	20	
NP_378194_467	33.1	2.80E+00	53	17	32		ZP_002188313513	32.3	4.70E+00	204	51	25	
NP_253000_629	33.1	2.80E+00	245	42	17		ZP_00171925503	32.3	4.70E+00	263	48	18	
YP_063897_576	33.1	2.80E+00	263	46	17		ZP_00201727822	32.3	4.70E+00	306	59	19	
YP_041192_424	33.1	2.80E+00	147	36	24		ZP_00149920505	32.3	4.70E+00	139	31	22	
NP_936490_622	33.1	2.80E+00	120	30	25		ZP_00133930158	32.3	4.70E+00	106	26	24	
NP_786249_1106	33.1	2.80E+00	330	64	19		ZP_00098108451	32.3	4.70E+00	139	33	23	
NP_777771_638	33.1	2.80E+00	121	29	23		ZP_00312864422	32	6.20E+00	142	36	25	
NP_764984_3692	33.1	2.80E+00	148	30	20		ZP_00298543733	32	6.20E+00	278	60	17	
NP_764984_3692	32.3	4.70E+00	264	52	19		ZP_003410411311	32	6.20E+00	230	37	16	
NP_600104_441	33.1	2.80E+00	280	63	22		ZP_00208461932	32	6.20E+00	211	43	20	
NP_693909_466	33.1	2.80E+00	126	24	19		ZP_00046132679	32	6.20E+00	143	32	22	
NP_608047_400	33.1	2.80E+00	108	29	26		NP_279901_636	32	6.20E+00	244	53	21	
NP_604026_1193	33.1	2.80E+00	253	52	20		NP_454751_1066	32	6.20E+00	277	56	20	
NP_559170_396	33.1	2.80E+00	119	29	24		NP_885621_1196	32	6.20E+00	65	18	27	
NP_541885_131	33.1	2.80E+00	62	24	38		AAQ60246_251	32	6.20E+00	98	30	30	

Fig. 25C-Continued

Fig. 25D

Hit_ID	Expected	Length	#Ident	%S	Hit_ID	Expected	Length	#Ident	%S
AADY01000C01	1.00E-100	373	213	57	spara_B_SPA_0.22632	4.00E-74	175	147	84
NZ_AADY01000001	1.00E-100	373	213	57	spara_B_SPA_0.22632	2.00E-37	100	82	82
NC_005126	1.00E-100	356	207	56	NZ_002928	3.00E-72	392	180	45
Yersinia	7.00E-89	364	202	55	NC_002928	1.00E-06	136	42	30
Yersinia	9.00E-69	367	200	54	NC_002929	6.00E-71	391	182	46
Yersinia	7.00E-59	308	149	48	NC_002929	1.00E-06	136	43	31
Yersinia	2.00E-31	294	107	36	NC_002927	2.00E-70	391	183	46
NC_005810	5.00E-86	368	202	54	NC_002927	1.00E-06	136	42	30
NC_005810	3.00E-21	223	57	30	NZ_AAAI02000003	1.00E-87	179	134	74
NC_005810	1.00E-04	66	22	33	NZ_AAAI02000003	6.00E-31	92	68	73
NC_003143	5.00E-86	368	202	54	AAAII02000003	1.00E-67	179	134	74
NC_003143	3.00E-21	223	57	30	AAAII02000003	6.00E-31	92	68	73
NC_003143	4.00E-19	295	76	25	NZ_AAAUJ02000002	9.00E-67	262	144	54
NC_003143	6.00E-18	309	79	25	NZ_AAAUJ02000002	4.00E-32	90	71	78
NC_006156	5.00E-86	368	202	54	AAAII02000002	9.00E-67	262	144	54
NC_008155	9.00E-20	207	61	29	AAAII02000002	4.00E-32	90	71	78
NC_008155	3.00E-08	115	37	32	AADX01000003	2.00E-65	297	145	48
NC_004088	5.00E-86	368	202	54	NZ_AADX01000003	2.00E-65	297	145	48
NC_004088	3.00E-21	223	67	30	NC_005085	7.00E-65	297	137	46
NC_004088	2.00E-07	84	28	33	NC_005085	3.00E-64	297	137	46
NC_002655	5.00E-82	274	169	61	NC_005085	1.00E-43	299	104	34
NC_002655	5.00E-33	115	74	64	NC_005085	9.00E-41	369	122	33
NC_002655	9.00E-06	299	66	22	NC_005085	6.00E-08	299	62	20
NC_002655	2.60E+00	185	42	22	NC_005085	6.80E-01	266	61	22
NC_002695	5.00E-82	274	169	61	NZ_AAAN02000069	9.00E-65	296	145	48
NC_002695	5.00E-33	115	74	64	NZ_AAAN02000069	5.00E-61	295	136	46
NC_002695	9.00E-06	299	66	22	NZ_AAAN02000069	6.00E-61	295	136	46
NC_002695	2.60E+00	185	42	22	NZ_AAAN02000069	1.00E-60	295	135	45
sentrifidis_716_10.21	5.00E-82	161	160	99	NZ_AAAN02000069	1.00E-59	296	132	44
shig277d05.q1k	5.00E-82	299	183	51	NZ_AAAN02000069	1.00E-59	296	132	44
shig277d06.q1k	6.00E-35	92	76	82	NZ_AAAN02000069	2.00E-46	296	112	37
Epath054a03.p1k	1.00E-80	268	171	64	AAAN02000069	9.00E-65	296	145	48
Epath054a03.p1k	1.00E-32	102	69	67	AAAN02000069	5.00E-61	295	138	46
NC_004431	3.00E-80	223	157	70	AAAN02000069	6.00E-61	295	136	46
NC_004431	6.00E-32	255	106	41	AAAN02000069	1.00E-60	295	135	45
Eagg66e09.q1k	2.00E-79	237	161	67	AAAN02000069	1.00E-59	296	132	44
Eagg66e09.q1k	9.00E-33	295	105	35	AAAN02000069	1.00E-59	296	132	44
NC_004337	1.00E-78	202	154	76	AAAN02000059	2.00E-46	296	112	37
NC_004337	1.00E-32	116	71	61	NZ_AAAN02000032	1.00E-62	296	134	45
NC_004741	1.00E-78	202	154	76	NZ_AAAN02000032	3.00E-61	296	133	44
NC_004741	1.00E-32	116	71	61	NZ_AAAN02000032	3.00E-61	296	133	44
dys055h06.q1k	2.00E-78	237	159	67	NZ_AAAN02000032	4.00E-60	296	136	45
dys055h06.q1k	2.00E-31	100	67	67	NZ_AAAN02000032	7.00E-56	295	130	44
NC_000913	1.00E-76	177	145	82	NZ_AAAN02000032	3.00E-53	282	117	41
NC_000913	4.00E-35	151	80	52	NZ_AAAN02000032	3.00E-17	87	47	54
NC_004757	5.00E-76	297	159	53	NZ_AAAN02000032	7.00E-12	275	64	23
NC_003198	8.00E-76	175	150	85	AAAN029600032	1.00E-62	296	134	45
NC_003198	4.00E-35	91	76	83	AAAN02000032	3.00E-61	296	133	44
NC_004631	8.00E-76	175	150	85	AAAN02000032	3.00E-61	296	133	44
NC_004631	4.00E-35	91	76	83	AAAN02000032	4.00E-59	296	136	45
NC_006511	1.00E-75	279	165	59	AAAN02000032	7.00E-56	295	130	44
NC_006511	4.00E-74	175	147	84	AAAN02000032	3.00E-53	282	117	41
NC_006511	2.00E-37	100	82	82	AAAN02000032	3.00E-17	87	47	54
NC_006511	5.00E-37	100	81	81	AAAN02000032	7.00E-12	275	64	23
NC_003197	1.00E-75	232	160	68	AAAT03000008	4.00E-62	297	138	46
NC_003197	9.00E-75	175	148	84	NZ_AAAT03000008	4.00E-62	297	138	46
NC_003197	5.00E-37	100	81	81	AADF01000011	2.00E-61	298	131	44
NC_003197	4.00E-36	98	79	80	AADF01000011	2.00E-10	322	67	20
spara_B_SPA_0.15635	1.00E-75	279	165	59	AADF01000011	3.10E-01	264	56	21
spara_B_SPA_0.15635	1.00E-31	74	55	87	NZ_AADF01000011	2.00E-61	295	131	44
spara_B_SPA_0.15635	1.00E-31	26	16	61	NZ_AADF01000011	2.00E-10	322	67	20
saft7-76a12.p1k	1.00E-75	232	160	68	NZ_AADF01000011	3.10E-01	264	56	21
saft7-76a12.p1k	5.00E-37	100	81	81	NC_004347	3.00E-61	297	140	47
NC_003916	3.00E-75	296	167	56	NC_004347	2.00E-60	297	138	46
saft11-469e08.p1k	9.00E-75	175	148	84	NC_004347	5.00E-07	136	36	26
saft11-469e08.p1k	4.00E-36	98	79	80	NC_004347	1.50E+00	89	26	29
bong455h05.q1k	3.00E-74	175	147	84	NZ_AAS02000027	8.00E-61	296	137	46
bong455h05.q1k	2.00E-36	100	80	80	AAAAS02000027	8.00E-61	296	137	46

Fig. 2SD-Continued

Hit_ID	Expected	Length	#ident	%S	Hit_ID	Expected	Length	#ident	%S
NC_006510	8.00E-51	307	134	43	J10154Af04.p1k	1.00E-51	300	115	38
NC_006510	4.00E-33	269	98	35	NC_004603	6.00E-51	376	131	34
NC_006510	3.00E-21	179	62	34	NC_004603	2.00E-42	384	125	32
NC_006510	2.00E-14	288	69	23	NC_004603	2.00E-40	377	129	33
config:3337:p_fluorescens	7.00E-60	300	134	44	NC_004603	2.00E-40	377	126	33
config:3337:p_fluorescens	5.00E-06	130	37	28	NC_004603	2.00E-40	377	126	33
NZ_AABP02000008	1.00E-59	297	135	45	NC_004603	3.00E-23	148	60	40
NZ_AABP02000008	4.00E-08	138	41	29	NC_004603	1.00E-12	217	61	28
AABP02000008	1.00E-59	297	135	45	NZ_AABG03000012	9.00E-51	295	120	40
AABP02000008	4.00E-08	138	41	29	NZ_AABG03000012	2.00E-48	292	118	40
AAE101000040	2.00E-59	291	134	46	NZ_AABG03000012	4.00E-20	274	89	25
NZ_AAE101000040	2.00E-59	291	134	46	AABG03000012	9.00E-51	295	120	40
bstearo.fasta.screen.Contig	4.00E-59	292	131	44	AABG03000012	2.00E-48	292	116	40
bstearo.fasta.screen.Contig	1.00E-54	327	133	40	AABG03000012	4.00E-20	274	69	25
bstearo.fasta.screen.Contig	1.00E-14	291	68	23	NZ_AAEK01000005	1.00E-50	309	119	38
NC_002939	6.00E-59	298	133	44	NZ_AAEK01000005	3.00E-49	293	117	38
NC_002939	2.00E-17	293	76	25	NZ_AAEK01000005	4.00E-49	292	118	40
config:4013:c_hydrogenoform	2.00E-58	298	133	44	NZ_AAEK01000005	1.00E-45	295	111	37
config:4013:c_hydrogenoform	8.00E-21	288	81	28	NZ_AAEK01000005	6.00E-32	292	96	32
NC_004578	1.00E-57	297	129	43	AAEK01000005	1.00E-50	309	119	38
NC_004578	1.50E+00	123	31	24	AAEK01000005	3.00E-49	293	117	38
NC_003869	1.00E-56	298	134	44	AAEK01000005	4.00E-49	292	118	40
NC_003869	9.00E-20	290	83	28	AAEK01000005	1.00E-45	295	111	37
NZ_AAAV02000005	3.00E-55	329	133	40	AAEK01000005	6.00E-32	292	98	32
AAAV02000005	3.00E-56	329	133	40	NC_005363	4.00E-50	295	124	42
NC_004344	3.00E-56	181	109	60	NC_005363	3.00E-49	292	120	41
NC_004344	2.00E-26	345	109	31	NC_005363	5.00E-47	301	117	38
NC_006270	5.00E-55	313	135	43	NC_005363	3.00E-46	292	115	39
NC_006270	2.00E+00	87	22	25	NC_005363	2.00E-43	295	110	37
NC_006322	5.00E-55	313	135	43	NC_005363	4.00E-42	295	109	36
NC_006322	2.00E+00	37	22	25	AAEM01000002	8.00E-60	404	145	36
Cbot440b12.q1c	6.00E-55	298	127	42	AAEM01000002	2.00E-48	404	150	37
Cbot440b12.q1c	1.00E-64	298	127	42	NZ_AAEM01000002	8.00E-60	404	146	36
Cbot440b12.q1c	4.00E-40	304	103	33	NZ_AAEM01000002	2.00E-48	404	150	37
Cbot440b12.q1c	2.00E-37	280	84	30	NC_003919	3.00E-49	212	113	53
Cbot440b12.q1c	3.00E-11	305	63	20	NC_003919	2.00E-21	307	67	28
NC_002570	1.00E-54	294	124	42	NC_003919	5.00E-09	134	40	39
NC_002570	2.00E-29	135	70	51	config:492:b_thailandensis	7.00E-49	380	136	35
NG_002570	4.00E-19	169	57	33	config:492:b_thailandensis	8.90E-01	49	18	36
Cd183h6.p1t	5.00E-54	300	125	41	sdubin_Config2945_12.23	9.00E-49	106	104	98
Cd183h6.p1t	3.00E-08	293	62	21	sdubin_Config2945_10.21	9.00E-49	106	104	98
Cd183h5.p1t	1.50E+00	309	55	17	NZ_AAFH01000001	2.00E-48	290	120	41
Cd183h5.p1t	2.00E+00	113	24	21	NZ_AAFH01000001	2.00E-24	159	67	42
AADW01000019	1.00E-53	299	128	42	AAFH01000001	2.00E-48	290	120	41
AADW01000019	7.00E-46	300	110	36	AAFH01000001	2.00E-24	159	67	42
AADW01000019	9.00E-12	321	74	23	NC_003902	2.00E-48	241	119	49
NZ_AADW01000019	1.00E-53	299	128	42	NC_003902	1.00E-21	319	93	29
NZ_AADW01000019	7.00E-46	300	110	36	NC_003902	1.00E-07	134	38	28
NZ_AADW01000019	9.00E-12	321	74	23	NC_006138	3.00E-43	252	120	47
NC_004557	2.00E-53	298	125	42	NC_006138	1.00E-43	231	108	46
NC_004557	1.00E-41	294	105	35	NC_006138	5.00E-25	227	81	35
NC_004557	3.00E-33	270	80	28	NC_006138	2.00E-22	254	79	31
NC_004557	1.00E-10	284	68	23	NC_006138	9.00E-08	130	40	30
NC_004557	2.00E+00	103	24	23	NC_006138	1.50E+00	146	33	22
NC_004193	2.00E-53	338	131	38	NC_005823	1.00E-47	302	111	36
NC_004193	1.00E-17	293	76	25	NC_005823	1.00E-43	300	107	35
BC10B6Le10.q1ka	2.00E-53	381	141	37	NC_005823	2.00E-43	300	107	35
BC10B5Le10.q1ka	4.00E-07	183	45	24	NC_005823	6.00E-39	301	99	32
NC_000964	6.00E-63	306	130	42	NC_005823	1.00E-08	116	34	30
NC_000964	1.00E-30	206	81	39	NC_005823	2	59	18	30
NC_004605	1.00E-82	301	119	39	NC_004342	1.00E-47	302	111	36
NC_004605	3.10E-01	116	28	25	NC_004342	1.00E-43	300	107	35
NC_003030	1.00E-52	295	122	41	NC_004342	3.00E-43	300	107	35
NC_003030	3.00E-45	292	115	39	NC_004342	5.00E-39	301	100	33
NC_003030	1.00E-41	290	104	35	NC_004342	1.00E-06	110	34	30
NC_003030	1.00E-31	292	89	30	NC_004342	2.00E+00	58	18	30
NC_003030	3.00E-08	289	65	22	NC_002505	2.00E-47	375	130	34
NC_003030	6.00E-03	310	69	22	NC_002505	5E-45	375	133	35

Fig. 25D-Continued

Hit_ID	Expected	Length	#Ident	%S	Hit_ID	Expected	Length	#Ident	%S
NC_002505	2.00E-43	377	134	36	NC_003296	4.00E-01	260	58	21
NC_002506	4.00E-43	377	129	34	NC_003296	5.20E-01	247	57	23
NC_002506	2.00E-41	379	127	33	NZ_AABN02000015	9.00E-43	300	105	36
NC_002506	2.00E-03	228	52	22	NZ_AABN02000015	2.00E-42	295	99	33
NC_006348	2.00E-47	385	135	35	AABN02000015	9.00E-43	300	108	36
NC_008348	1.50E+00	78	24	31	AABN02000015	2.00E-42	295	99	33
Burkholderia	2.00E-47	385	136	35	NZ_AADQ01000005	2.00E-42	298	107	36
Burkholderia	2.00E-02	336	74	22	NZ_AADR01000002	2.00E-42	298	107	36
Burkholderia	1.50E+00	76	24	31	AADR01000002	2.00E-42	296	107	36
NC_004722	3.00E-47	296	114	38	AADQ01000005	2.00E-42	298	107	36
NC_004722	2.00E-44	298	115	38	NC_003212	2.00E-42	296	107	36
NC_004722	6.00E-40	271	98	36	NC_003212	6.20E-02	261	57	21
NC_004722	2.00E-32	291	95	32	NC_003212	1.40E-01	292	54	21
NC_004722	3.00E-07	243	57	23	NC_003210	2.00E-42	296	107	36
NC_008177	4.00E-47	295	114	38	NC_003210	6.00E-03	273	58	21
NZ_AAECR01050023	6.00E-47	292	112	38	NC_002973	2.00E-42	298	107	36
NZ_AAE01000034	6.00E-47	292	112	38	NC_005139	2.00E-42	375	121	32
NZ_AAE01000025	6.00E-47	292	112	38	NC_006139	2.00E-36	376	118	31
NZ_AAE01000011	6.00E-47	292	112	38	NC_005139	9.00E-36	376	119	31
NZ_AAEF01000035	6.00E-47	292	112	38	NC_005139	4.00E-35	178	80	44
NZ_AAEQ01000029	6.00E-47	292	112	38	NC_005139	7.00E-35	376	111	29
AAES01000034	6.00E-47	292	112	38	NC_005139	7.00E-37	374	108	28
AAER01000023	6.00E-47	292	112	38	NC_005139	6.00E-21	301	92	30
AAEQD01000029	6.00E-47	292	112	38	NC_004459	2.00E-42	375	121	32
AAEP01000035	6.00E-47	292	112	38	NC_004459	2.00E-41	378	122	32
AAEO01000025	6.00E-47	292	112	38	NC_004459	7.00E-36	378	120	31
AAEN01000011	6.00E-47	292	112	38	NC_004459	9.00E-36	376	119	31
NC_009845	6.00E-47	292	112	38	NC_004459	4.00E-35	178	80	44
NC_007530	6.00E-47	292	112	38	NC_004459	7.00E-27	374	108	28
NC_003597	6.00E-47	292	112	38	NC_004459	6.00E-21	301	92	30
NC_003985	6.00E-47	292	112	38	NC_006187	4.00E-42	299	105	35
NC_002367	1.00E-46	300	113	37	NC_005087	3.00E-20	308	85	27
NC_002967	1.00E-43	304	109	35	NC_008613	6.00E-42	161	89	58
NC_002967	5.00E-42	304	106	34	Eagg631c06.q1k	8.00E-42	310	107	34
NC_002967	3.00E-10	285	66	22	Eagg631c06.c1k	8.00E-08	309	62	20
NC_006370	9.00E-46	382	130	34	NZ_AAJ03000013	1.00E-41	378	127	33
NC_006370	9.00E-36	251	106	42	AAAJ03000013	1.00E-41	378	127	33
NC_006370	5.00E-34	213	82	38	NZ_AABQ07000002	1.00E-40	242	103	42
NC_006370	2.00E-17	203	63	31	NZ_AABQ07000002	3.00E-24	335	100	29
NC_006370	4.00E-17	149	55	36	NZ_AABQ07000002	3.00E-04	134	34	25
NC_006370	2.00E+00	73	16	21	AABQ07000002	1.00E-40	242	103	42
NC_000919	2.00E-45	298	114	38	AABQ07000002	3.00E-24	335	100	29
NC_000919	1.00E-44	301	111	36	AABQ07000002	3.00E-04	134	34	25
NC_000919	1.00E-44	304	107	35	NC_002516	1.00E-40	242	103	42
NC_000919	3.00E-09	268	66	22	NC_002516	3.00E-24	335	100	29
NC_002937	3.00E-45	311	119	38	NC_002516	3.00E-04	134	34	25
NC_002937	1.00E-43	310	113	36	NC_002516	3.10E-01	201	49	24
NC_002937	2.00E-41	295	101	34	AABE03000002	7.00E-10	223	101	45
NC_002937	2.00E+00	171	36	21	AABE03000002	5.00E-39	254	105	41
NZ_AABN02000006	6.00E-46	295	107	36	AABE03000002	4.00E-20	182	66	35
NZ_AABN02000006	3.00E-03	46	21	45	AABE03000002	4.00E-18	129	55	42
AABN02000006	6.00E-45	295	107	36	AABE03000002	1.00E-06	128	33	25
AABN02000006	3.00E-03	46	21	45	AABE03000002	2.80E-02	91	24	26
scublin_Config1652_12.23	1.00E-44	97	97	100	AABE03000002	1.40E-01	169	35	20
NC_006368	2.00E-44	309	124	40	NZ_AABE03000002	7.00E-40	222	101	45
NC_006368	1.00E-23	200	77	38	NZ_AABE03000002	5.00E-39	254	105	41
NC_005957	3.00E-44	365	125	34	NZ_AABE03000002	4.00E-20	182	66	36
NC_005957	5.00E-33	292	97	33	NZ_AABE03000002	4.00E-19	128	55	42
NC_002942	5.00E-44	309	126	40	NZ_AABE03000002	1.00E-08	128	33	25
NC_002942	3.00E-23	198	78	39	NZ_AABE03000002	2.80E-02	91	24	26
NC_005369	2.00E-43	309	122	39	NZ_AABE03000002	1.40E-01	169	35	20
NC_006369	2.00E-23	251	88	35	config:1731:c_psychroerythr	7.00E-40	269	108	41
NC_006156	3.00E-43	342	113	33	config:1731:c_psychroerythr	6.00E-39	259	108	41
NC_006155	8.90E-01	77	18	23	config:1731:c_psychroerythr	6.00E-24	263	82	31
NC_001318	3.00E-43	342	113	33	config:1731:c_psychroerythr	1.00E-21	263	78	28
NC_001318	8.90E-01	77	18	23	config:1731:c_psychroerythr	8.00E-07	135	32	23
NC_003296	4.00E-43	297	109	36	NC_002947	1.00E-39	252	103	40
NC_003296	7.00E-03	254	60	22	NC_002947	3.00E-19	128	53	41

Fig. 25D-Continued

Hit_ID	Expected	Length	#Ident	%S	Hit_ID	Expected	Length	#Ident	%S
NC_002947	4.00E-07	130	41	31	NC_00853	8.00E-10	256	62	24
AAAW02000007	2.00E-39	385	128	33	pputidprs_GPPBG16TR	7.00E-28	123	66	53
AAAW02000007	8.00E-23	298	89	29	std125b05_p1k	9.00E-25	77	58	75
AAAW02000007	2.00E-09	154	47	30	AAEF01000080	3.00E-24	245	76	31
NZ_AAAW02000007	2.00E-39	385	128	33	AAEF01000080	7.00E-20	195	65	33
NZ_AAAW02000007	8.00E-23	298	89	29	AAEF01000080	3.00E-13	301	76	25
NZ_AAAW02000007	2.00E-09	154	47	30	NZ_AAEF01000080	3.00E-24	245	76	31
NC_004463	1.00E-38	295	95	32	NZ_AAEF01000080	7.00E-20	195	65	33
NC_004463	8.00E-16	311	73	23	NZ_AAEF01000080	3.00E-13	301	76	25
NC_004463	1.00E-14	317	74	23	NC_005090	3.00E-23	294	89	30
NC_004463	8.00E-08	273	72	26	NC_005090	2.00E-21	192	63	32
NC_004463	2.00E-07	255	72	26	NC_005090	9.00E-19	333	85	25
NC_004463	8.00E-07	250	63	25	NC_005090	1.00E-12	278	65	23
NC_004463	2.00E-06	186	47	25	NC_005090	3.00E-12	122	44	36
NC_004463	8.10E-02	233	50	21	NC_005090	6.00E-03	168	37	22
NC_004463	1.40E-01	134	39	29	config:521:c_jejuni	3.00E-23	181	66	36
NC_004463	4.00E-01	109	29	26	config:521:c_jejuni	1.00E-21	181	66	36
NC_004463	6.80E-01	121	28	23	config:521:c_jejuni	2.00E-14	156	53	33
NC_004463	1.20E+00	109	29	26	config:521:c_jejuni	2.00E-12	156	50	32
NC_004463	1.50E+00	109	29	26	config:521:c_jejuni	6.00E-11	288	68	23
NC_004463	2.00E+00	109	29	26	NC_00915	8.00E-23	187	67	35
Pflu019c02.p1c	2.00E-38	230	97	42	NC_00915	3.00E-19	170	57	33
Pflu019c02.p1c	2.00E-17	330	81	24	NC_00915	5.00E-15	311	78	25
contig:491:b_thailandensis	3.00E-38	305	107	35	NC_00915	2.00E-13	121	47	38
AAEH01000086	2.00E-37	242	97	40	NC_00921	1.00E-22	187	67	35
AAEH01000086	2.00E-17	86	46	53	NC_00921	3.00E-19	170	57	33
NZ_AAEH01000086	2.00E-37	242	97	40	NC_00921	5.00E-15	311	78	25
NZ_AAEH01000086	2.00E-17	86	46	53	NC_00921	2.00E-13	121	47	38
AAAG02000001	1.00E-36	305	107	35	NC_00921	4.00E-04	149	40	25
AAAG02000001	4.00E-12	302	69	22	NC_002163	3.00E-22	171	63	36
AAAG02000001	6.80E-01	239	49	20	NC_002163	6.00E-21	181	66	36
NZ_AAAG02000001	1.00E-36	305	107	35	NC_002163	4.00E-14	132	48	36
NZ_AAAG02000001	4.00E-12	302	69	22	NC_002163	4.00E-12	132	45	34
NZ_AAAG02000001	6.80E-01	239	49	20	NC_002163	7.00E-11	288	66	22
NC_005296	4.00E-36	295	92	31	NC_004917	9.00E-22	188	67	35
NC_005296	2.00E-05	284	70	24	NC_004917	1.00E-19	170	60	35
NC_005296	2.00E-05	284	69	24	NC_004917	1.00E-19	170	60	35
NC_005296	1.00E-03	194	49	25	NC_004917	3.00E-13	297	74	24
NC_005296	8.90E-01	261	58	22	NC_004917	3.00E-13	297	74	24
sdbifn_Config993_12.23	4.00E-36	81	81	100	NC_004917	6.00E-11	115	40	34
stdt38a03.p1k	4.00E-36	98	79	80	NC_002696	1.00E-21	297	76	25
stdt23f04.p1k	4.00E-36	98	79	80	NC_002696	4.00E-20	298	76	25
NZ_AADT02000011	2.00E-35	382	125	32	NC_002696	5.00E-20	292	75	25
NZ_AADT02000011	8.00E-18	285	73	25	NC_002696	5.00E-20	292	72	24
AADT02000011	2.00E-35	382	125	32	NC_002696	7.00E-20	292	72	24
AADT02000011	8.00E-18	285	73	25	NC_002696	1.00E-19	292	70	23
NC_006512	6.00E-35	170	82	48	NC_002696	1.10E-01	73	21	28
NC_006512	4.00E-20	267	77	28	NC_00918	1.00E-20	209	82	29
NC_006512	1.00E-03	130	35	26	NC_00918	7.00E-09	129	39	30
NZ_AAAI02000004	3.00E-34	199	85	42	config:4798:g_obscuriglobus	7.00E-20	139	57	41
NZ_AAAI02000004	9.00E-22	92	57	61	config:4798:g_obscuriglobus	3.00E-12	157	52	33
AAAII02000004	3.00E-34	199	85	42	AAED01000008	9.00E-20	294	80	27
AAAII02000004	9.00E-22	92	57	61	AAED01000008	8.00E-15	297	68	22
NC_003909	5.00E-34	292	96	32	NZ_AAE01000008	9.00E-20	294	80	27
NC_003909	2.00E-28	221	77	34	NZ_AAE01000008	8.00E-15	297	68	22
NC_003909	8.00E-13	84	41	48	NC_005027	9.00E-20	284	88	30
NZ_AAN02000082	5.00E-33	280	85	30	NC_005027	1.00E-19	257	76	29
AAAN02000082	5.00E-33	280	85	30	NC_005027	4.00E-18	169	62	36
NZ_AAAE01000066	1.00E-32	166	74	44	NC_005027	6.00E-16	126	55	43
NZ_AAAE01000066	2.00E-19	176	63	35	NC_005027	1.00E-14	143	57	39
AAAE01000066	1.00E-32	166	74	44	NC_005027	4.00E-14	250	70	28
AAAE01000066	2.00E-19	176	63	35	NC_005027	5.80E-01	240	52	21
NC_005274	1.00E-32	292	95	32	NC_005027	1.50E+00	120	34	28
NC_006274	1.00E-31	289	96	33	rhi2673a05.p1n	2.00E-19	303	84	27
NC_006274	1.00E-14	111	46	41	AAAW02000158	2.00E-18	314	89	28
NC_006274	1.00E-07	242	57	23	NZ_AAAI02000158	2.00E-18	314	89	28
NC_000853	2.00E-32	392	119	30	NC_003082	1E-17	313	77	24
NC_000853	3.00E-17	248	63	25	NC_003062	3.00E-14	313	75	23

Fig. 25D-Continued

Hit_ID	Expected	Length	#ident	%S	Hit_ID	Expected	Length	#ident	%S
NC_003062	9.00E-12	324	75	23	AAEA01000001	1.00E-03	295	62	21
NC_003062	2.00E-06	256	60	23	NZ_AAEAD1000001	1.00E-03	295	62	21
NC_003062	4.00E-04	140	34	24	NC_002662	1.00E-03	287	50	17
NC_003304	1.00E-17	313	77	24	Pflu562a01.p1kz	3.00E-03	277	59	24
NC_003304	3.00E-14	313	75	23	AABK03000008	3.00E-03	273	63	23
NC_003304	9.00E-12	324	75	23	NZ_AABK03000008	3.00E-03	273	63	23
NC_003304	2.00E-08	256	60	23	AAEHG1000003	3.00E-03	284	62	21
NC_003304	4.00E-04	140	34	24	AAEHG1000003	2.10E-02	282	56	19
NC_004311	7.00E-17	298	76	25	AAEHG1000003	2.80E-02	287	60	20
NC_003318	3.00E-16	298	75	25	AAEHG1000003	2.80E-02	295	59	20
sdublin_Config1018_12.23	3.00E-18	80	49	21	AAEHG1000003	3.80E-02	285	61	21
rhlz606c07.p1n	4.00E-16	303	78	25	AAEHG1000003	4.70E-02	285	61	21
rhlz606c07.p1n	5.00E-15	304	76	25	AAEHG1000003	8.10E-02	285	69	20
rhlz606c07.p1n	1.00E-05	317	66	20	AAEHG1000003	1.10E-01	285	60	21
NC_002678	1.00E-15	329	75	22	AAEHG1000003	1.10E-01	322	67	20
NC_002678	3.00E-08	356	80	22	NZ_AAEHG1000003	3.00E-03	264	62	21
NZ_AAFG01000002	2.00E-15	296	75	25	NZ_AAEHG1000003	2.10E-02	282	56	19
NZ_AAFG01000002	4.00E-14	298	72	24	NZ_AAEHG1000003	2.80E-02	287	60	20
AAFG01000002	2.00E-15	296	75	25	NZ_AAEHG1000003	2.80E-02	295	59	20
AAFG01000002	4.00E-14	298	72	24	NZ_AAEHG1000003	3.60E-02	285	61	21
rhlz24d12.p1k	3.00E-14	310	77	24	NZ_AAEHG1000003	4.70E-02	285	61	21
NZ_AAAE01000155	7.00E-14	302	70	23	NZ_AAEHG1000003	8.10E-02	285	59	20
AAAE01000155	7.00E-14	302	70	23	NZ_AAEHG1000003	1.10E-01	285	60	21
NZ_AAFG01000004	2.00E-13	294	72	24	NZ_AAEHG1000003	1.10E-01	322	67	20
AAFG01000004	2.00E-13	294	72	24	NC_005140	3.00E-03	289	63	21
NZ_AAFG01000007	3.00E-13	299	69	23	NZ_AADQ01000014	4.00E-03	258	56	21
AAFG01000007	3.00E-13	299	69	23	AADQ01000014	4.00E-03	258	56	21
config:519:c_ijeunii	6.00E-11	288	68	23	AAEI01000009	4.00E-03	304	61	20
NZ_AAAPC01001822	1.00E-10	276	72	26	AAEI01000009	4.00E-03	298	64	21
AAAP01001822	1.00E-10	276	72	26	AAEI01000009	8.10E-02	292	62	21
CboI389c10.p2c412	1.00E-10	257	63	24	AAEI01000009	1.80E-01	278	57	20
NC_003047	5.00E-09	321	68	21	NZ_AAEI01000009	4.00E-03	304	61	20
NC_003047	2.00E-05	87	29	33	NZ_AAEI01000009	4.00E-03	298	64	21
NC_003047	5.00E-04	245	56	23	NZ_AAEI01000009	8.10E-02	292	62	21
NC_003047	2.80E-02	87	24	27	NZ_AAEI01000009	1.80E-01	278	57	20
NZ_AAEK01000008	2.00E-08	243	62	25	NC_002935	6.00E-03	169	38	22
AAEK01000008	2.00E-08	243	62	25	NC_004567	7.00E-03	285	64	22
NZ_AABN02000001	3.00E-07	113	32	28	NC_004567	2.00E+00	299	58	19
NZ_AABN02000001	3.00E-03	238	50	21	NZ_AAUU02000026	1.00E-02	122	36	29
AABN02000001	3.00E-07	113	32	28	AAAU02000026	1.00E-02	122	36	29
AABN02000001	3.00E-03	238	50	21	AAEI01000011	1.20E-02	237	72	24
config:2663:s_pomeroyi	4.00E-07	278	64	23	NZ_AAEI01000011	1.20E-02	297	72	24
config:2663:s_pomeroyi	4.00E-03	180	36	29	NC_002758	1.20E-02	332	72	21
config:1062:h_nephrium	6.00E-07	163	39	25	NC_002746	1.20E-02	332	72	21
NZ_AAAI02000005	5.00E-06	326	70	21	NZ_AABH02000009	1.60E-02	241	60	24
NZ_AAAI02000005	2.00E+00	93	28	30	AABH02000009	1.60E-02	241	60	24
AAAI02000005	3.00E-06	326	70	21	NZ_AAAU02000005	2.10E-02	292	54	18
AAAI02000005	2.00E+00	93	28	30	AAAR02000005	2.10E-02	292	54	18
Bcep1110a03.q2kb4087	3.00E-05	288	66	22	NC_006449	2.80E-02	302	54	21
Bcep1110a03.q2kb4087	5.00E-05	299	67	22	NC_006448	2.80E-02	302	64	21
Bcep1110a03.q2kb4087	3.00E-04	286	66	23	sdublin_Config4745_12.23	2.80E-02	19	18	94
Bcep1110a03.q2kb4087	4.00E-01	226	52	23	sdublin_Config3450_12.23	2.80E-02	19	18	94
AAAY01000003	8.00E-06	121	37	30	mar499b01.q1k	3.60E-02	87	25	28
NZ_AADY01000003	8.00E-05	121	37	30	NZ_AAM01000127	4.70E-02	57	21	36
NZ_AABG03000053	2.00E-04	83	28	33	NZ_AACK01000006	4.70E-02	146	33	26
AAEI01000016	2.00E-04	291	67	23	AACK01000006	4.70E-02	146	38	26
AAEI01000016	9.00E-04	289	59	20	AAAM01000127	4.70E-02	57	21	36
AABG03000053	2.00E-04	83	28	33	NC_003923	4.70E-02	283	68	23
NZ_AAEI01000016	2.00E-04	291	67	23	NC_003923	1.60E+00	299	60	20
NZ_AAEI01000016	9.00E-04	289	59	20	NC_002953	4.70E-02	283	55	23
NZ_AABG03000001	4.00E-04	257	54	21	ap5.fasta.screen.Contig207	4.70E-02	145	38	26
AABG03000001	4.00E-04	257	54	21	NZ_AAN02000006	6.20E-02	294	57	19
config:313:m_arthritidis	4.00E-04	263	60	22	NZ_AAJJ03000006	6.20E-02	310	62	20
config:313:m_arthritidis	1.40E-01	127	32	NZ_AAJJ03000006	1.10E-01	308	61	19	
config:313:m_arthritidis	1.80E-01	124	33	NZ_AAJJ03000006	1.80E-01	272	54	19	
config:313:m_arthritidis	1.50E+00	137	32	NZ_AAJJ03000006	3.10E-01	275	55	20	
BcepB25a04.p2n33	4.00E-04	255	58	22	NZ_AAJJ03000006	3.10E-01	254	63	20
config:510:v_spinosum	7.00E-04	262	66	26	NZ_AAJJ03000006	6.80E-01	279	55	19

Fig. 25D-Continued

Hit_ID	Expected	Length	#Ident	%S	Hit_ID	Expected	Length	#Ident	%S
NZ_AAAJ03000006	8.90E-01	319	59	18					
NZ_AAAJ03000006	1.50E+00	324	59	18					
AAAN02000006	6.20E-02	294	57	19					
AAAJO3000006	6.20E-02	310	62	20					
AAAJO3000006	1.10E-01	308	61	19					
AAAJO3000006	1.80E-01	272	54	19					
AAAJO3000006	3.10E-01	275	56	20					
AAAJO3000006	3.10E-01	254	53	20					
AAAJO3000006	6.80E-01	279	55	19					
AAAJO3000006	8.90E-01	319	59	18					
AAAJO3000006	1.50E+00	324	59	18					
NZ_AAP01003482	8.10E-02	278	63	22					
AAAP01003482	8.10E-02	278	63	22					
AADW01000010	8.10E-02	317	71	22					
NZ_AADW01000010	8.10E-02	317	71	22					
NC_004461	8.10E-02	169	43	25					
NC_004461	1.20E+00	302	60	19					
config:370:s_epidermidis	8.10E-02	169	43	25					
contig:370:s_epidermidis	1.20E+00	302	60	19					
AAED01000003	1.40E-01	183	41	22					
NZ_AAED01000003	1.40E-01	183	41	22					
NZ_AAAN02000003	2.30E-01	260	53	20					
AAAN02000003	2.30E-01	260	53	20					
AADW01000069	2.30E-01	285	51	17					
NZ_AADW01000069	2.30E-01	285	51	17					
NZ_AABH02000010	2.30E-01	298	58	19					
AABH02000010	2.30E-01	298	58	19					
Bcep1157h02_p1c	3.10E-01	300	58	19					
NZ_AABG03000015	4.00E-01	181	39	21					
NZ_AAAN02000029	4.00E-01	89	26	29					
AAAN02000029	4.00E-01	89	26	29					
AABG03000015	4.00E-01	181	39	21					
NZ_AABQ07000001	4.00E-01	201	49	24					
AABQ07000001	4.00E-01	201	49	24					
NC_006300	4.00E-01	273	58	21					
NC_006300	4.00E-01	273	58	21					
NC_006300	5.20E-01	274	61	22					
NC_003272	5.20E-01	286	59	20					
NC_004829	5.20E-01	183	40	21					
NZ_AAAN02000076	6.80E-01	294	56	19					
AAAN02000076	6.80E-01	294	56	19					
NZ_AAFG01000003	8.90E-01	62	20	32					
AAFG01000003	8.90E-01	62	20	32					
NZ_AABM02000032	8.90E-01	165	39	23					
AABM02000032	8.90E-01	165	39	23					
NC_004663	8.90E-01	160	44	29					
contig:3563:m_smegmatis	8.90E-01	102	30	29					
AAE101000026	1.20E+00	130	38	29					
NZ_AAE101000026	1.20E+00	130	38	29					
NC_005125	1.20E+00	73	24	32					
NC_003366	1.20E+00	232	49	21					
NC_000951	1.20E+00	114	30	26					
Bcep115d11.q1k	1.20E+00	130	37	28					
contig:4304:c_perfringens	1.20E+00	232	49	21					
NZ_AAEV01000005	1.50E+00	187	41	21					
AAEV01000005	1.50E+00	187	41	21					
NZ_AABP02000001	1.50E+00	254	46	18					
AABP02000001	1.50E+00	254	46	18					
NC_003552	1.50E+00	257	49	19					
NC_005362	2.00E+00	125	28	22					
AAAY02000105	2.60E+00	112	29	25					
AAEH01000017	2.60E+00	238	54	22					
NZ_AAEH01000017	2.60E+00	238	54	22					
NZ_AAAY02000105	2.60E+00	112	29	25					
Eagg385b08.q1ka	2.60E+00	185	42	22					

Fig. 26

Accession	Source	Amino % Identity	Carboxy % Identity	Length
Q53970	Phase-I flagellin [fliC] [Salmonella dublin]	100	100	505
P72151	B-type flagellin [fliC] [Pseudomonas aeruginosa]	50	58	488
Q5X5M6	Flagelline [flaA] [Legionella pneumophila str. Paris]	46	61	475
Q6VMV6	Flagellin [fliC] [Escherichia coli]	74	84	488
P13713	Flagellin [fliC] [Serratia marcescens]	65	75	351
Q93RK8	Flagellin [tag] [Bacillus subtilis]	50	49	333
Q02551	Flagellin [flaA] [Listeria monocytogenes]	39	38	287
Q09012	Flagellin [fliC-SS(a)] [Shigella sonnei]	72	84	524
Q8GNT8	Flagellin [Edwardsiella tarda]	65	77	416
Q9FAE7	Flagellin [H8301-fla] [Acidovorax avenae subsp. avenae]	61	56	492
Q8ZF76	Thermoregulated motility protein (Putative flagellin) [fliC] [Yersinia pestis]	82	80	369
Q7N5I4	Flagellin [fliC] [Photobacterium luminescens (subsp. laumontii)]	63	77	355
Q33578	Flagellar filament protein [fliC] [Rhodobacter sphaeroides (Rhodopseudomonas sphaerooides)]	46	55	493
Q56826	Flagellin [fliC] [Xenorhabdus nematophilus]	76	75	313
P42273	Flagellin 2 [fliC2] [Proteus mirabilis]	63	78	367
O31059	Flagellin A protein [flaA] [Buvyrivibrio fibrisolvens]	53	56	472
Q7VZC2	Flagellin [flaA] [Bordetella pertussis]	74	77	391
Q9F4A4	Flagellin protein FlA(C) [fliC] [Clostridium chauvoei]	51	48	413
Q8PP9C4	Flagellar protein [fliC] [Xanthomonas campestris (pv. campestris)]	48	55	399
Q82UA3	Hag; flagellin [tag] [Nitrosomonas europaea]	58	57	275
Q84IC5	Flagellin (Fragment) [flaA] [Campylobacter lari]	32	45	490

Fig. 27

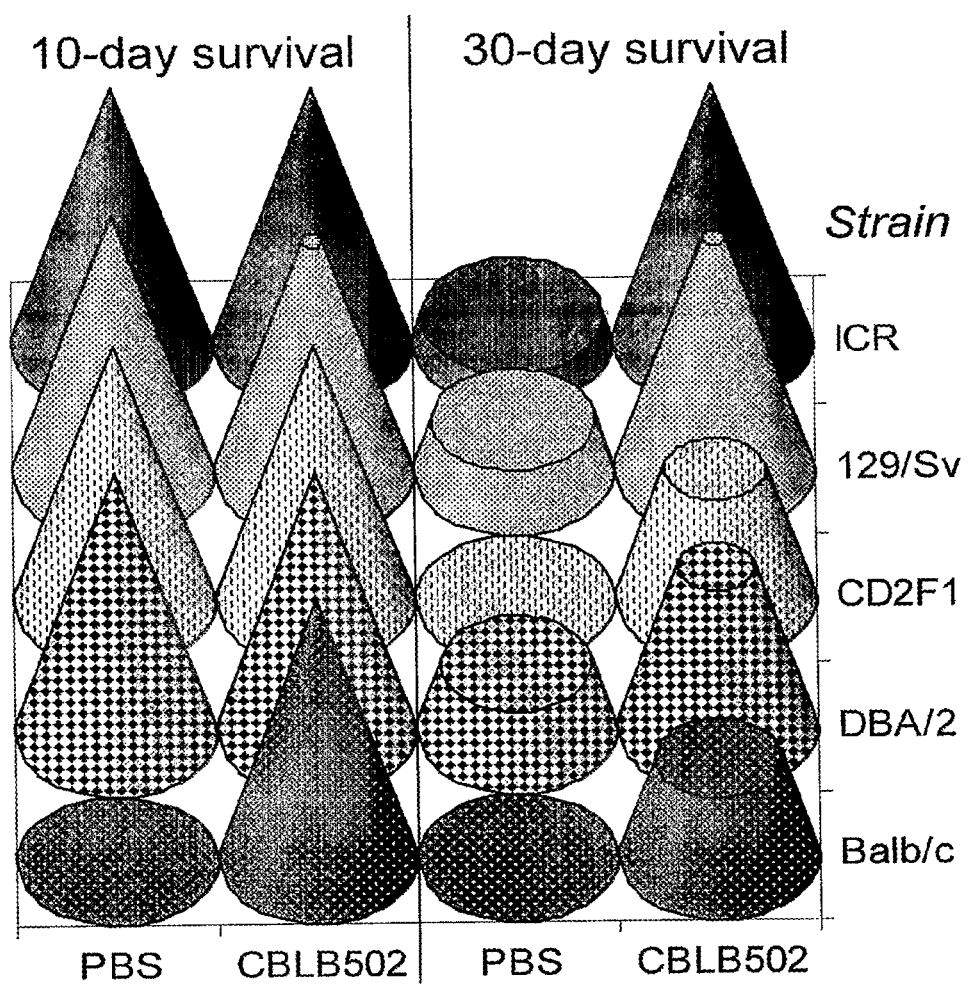


Fig. 28

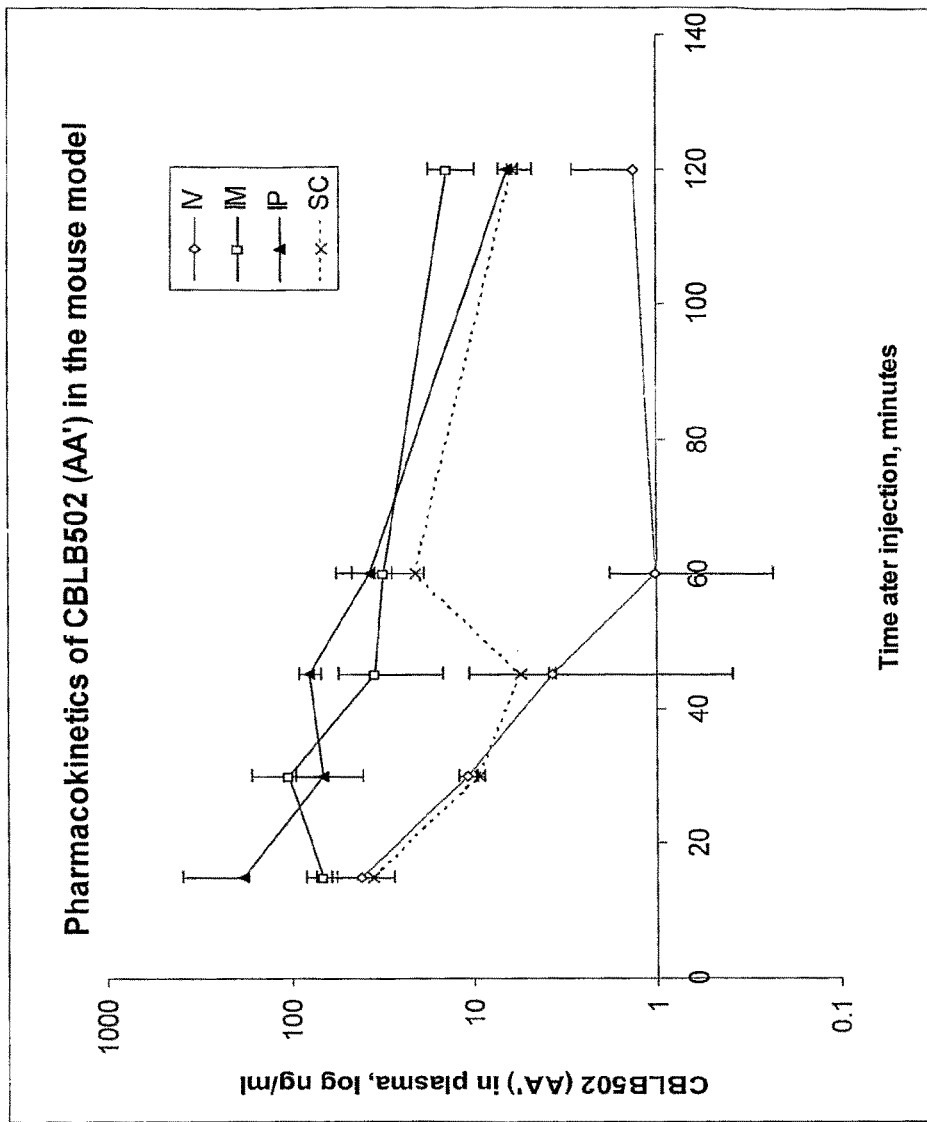


Fig. 29

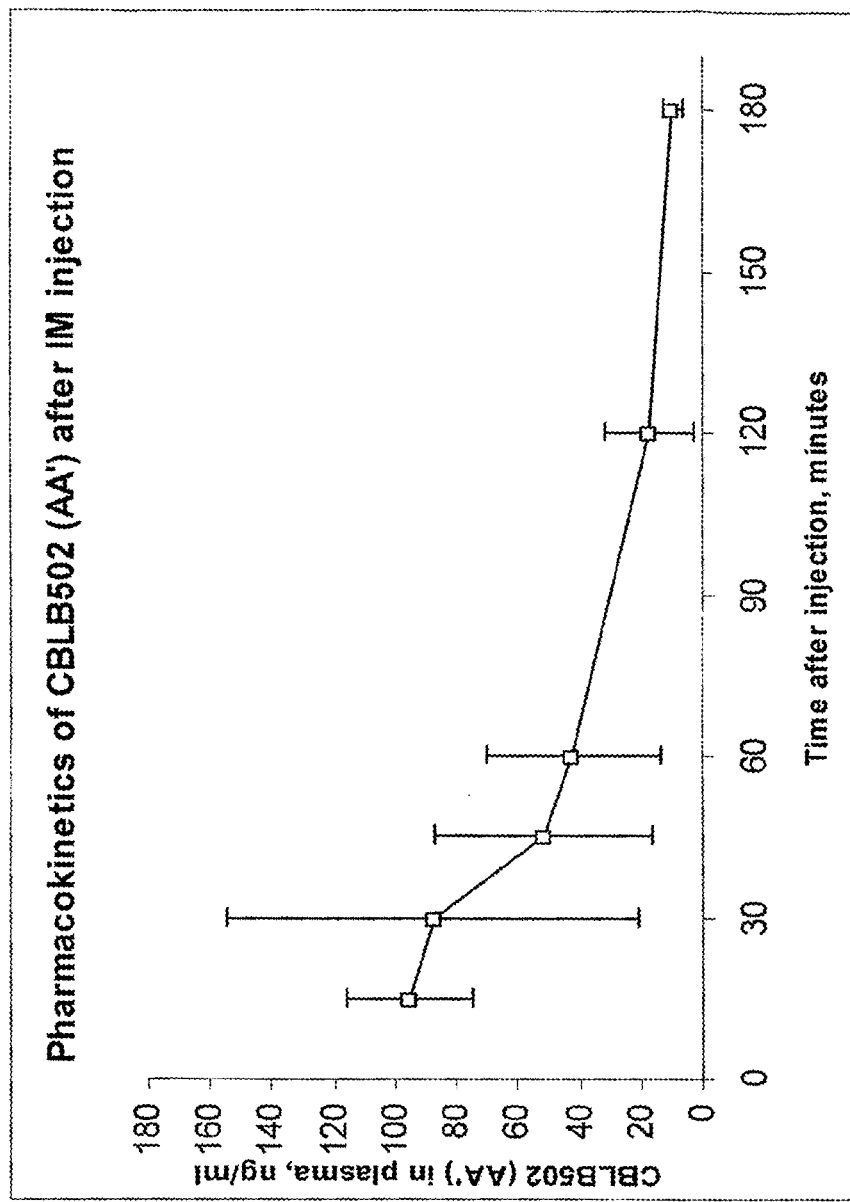


Fig. 30

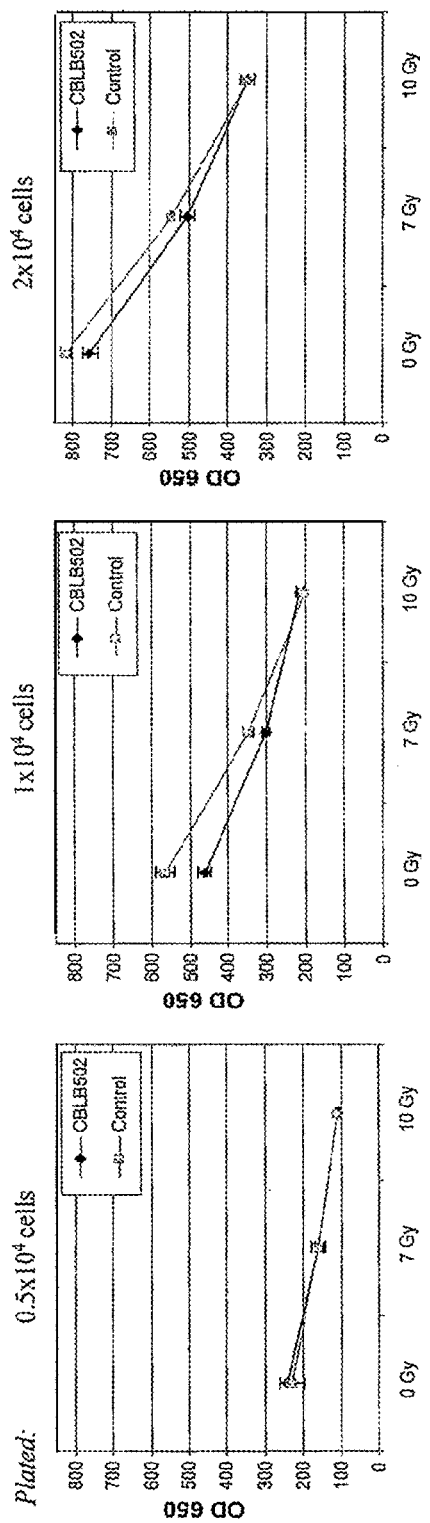


Fig. 31

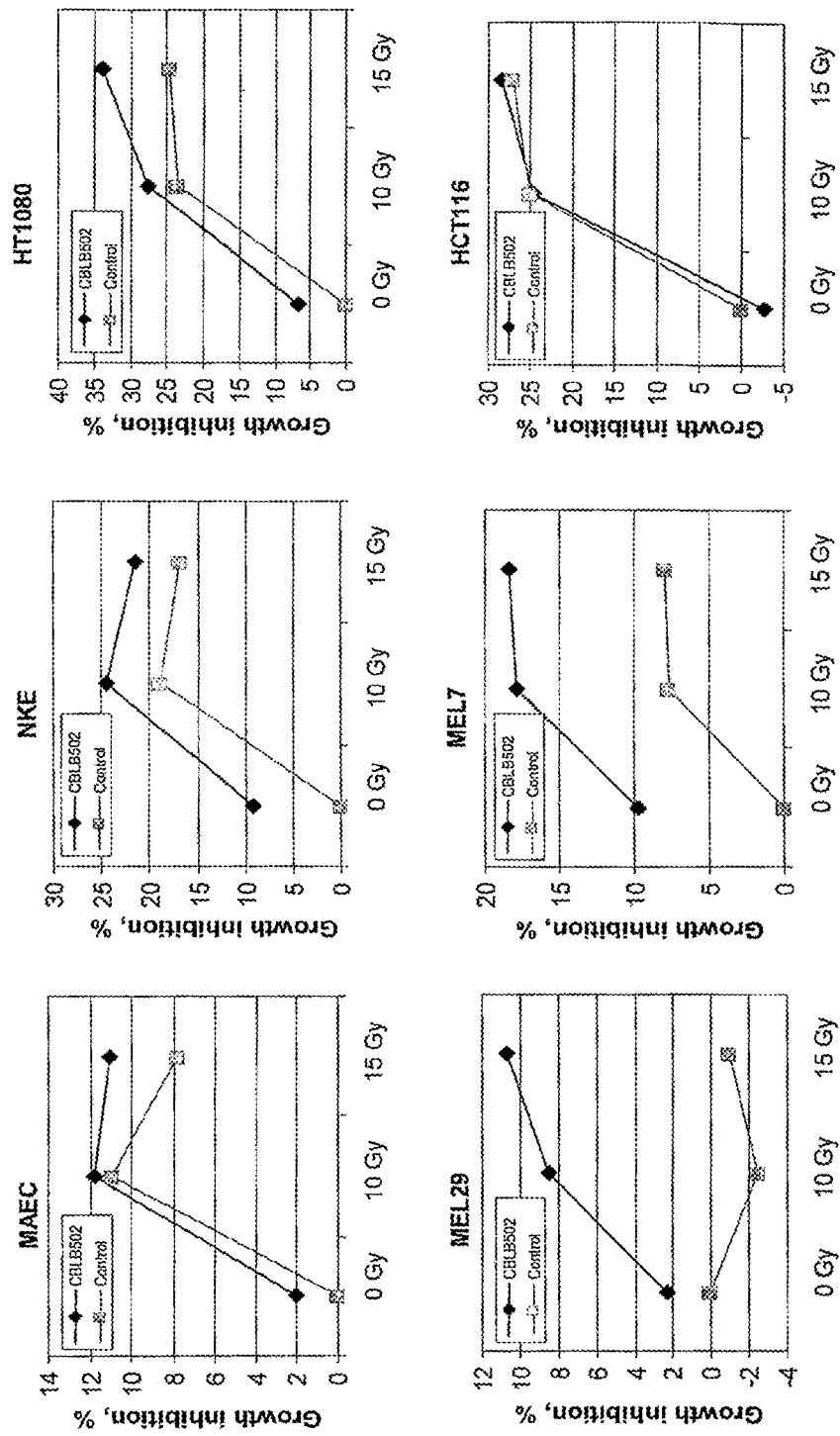


Fig. 32

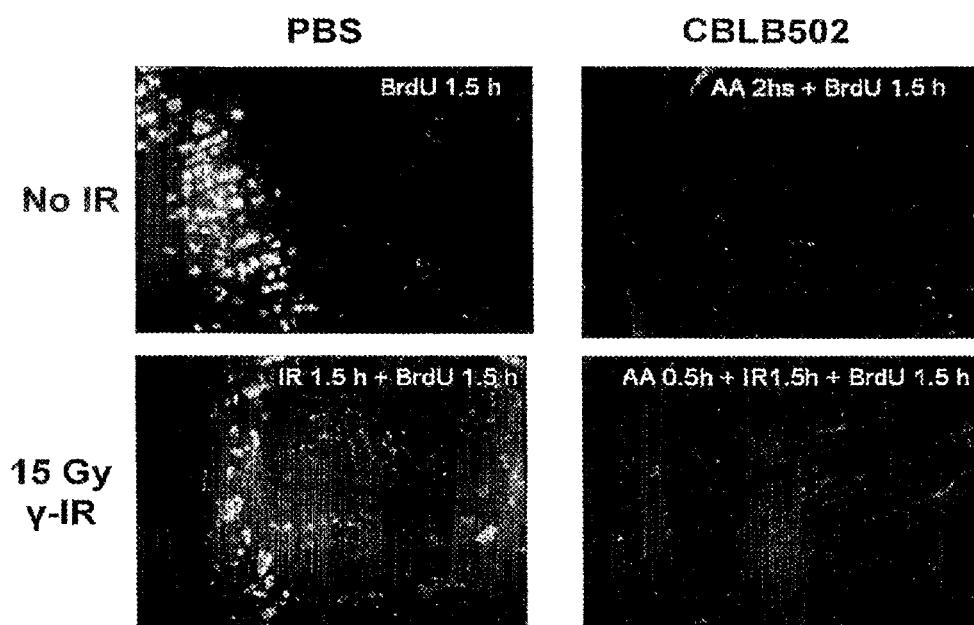


Fig. 33

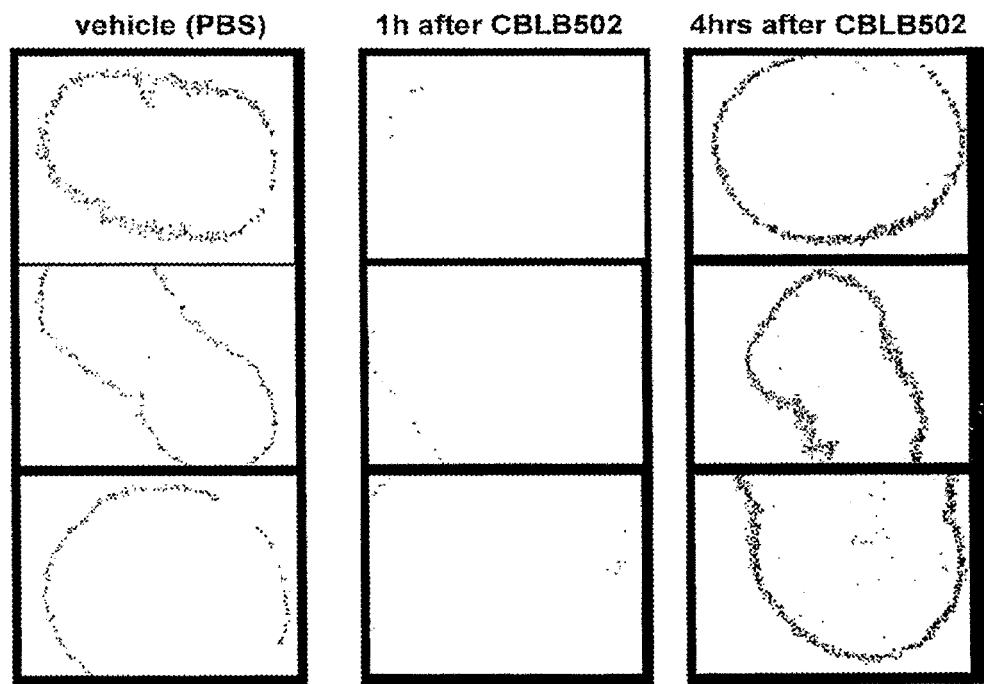


Fig. 34

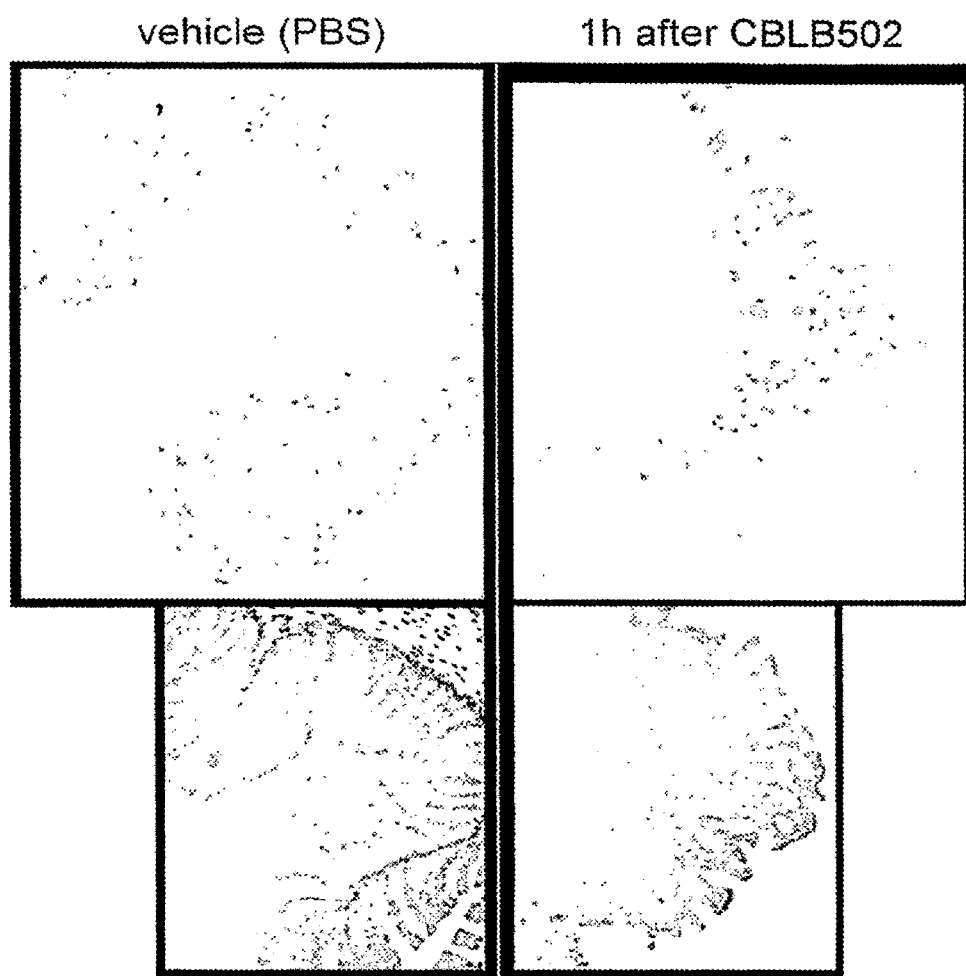


Fig. 35

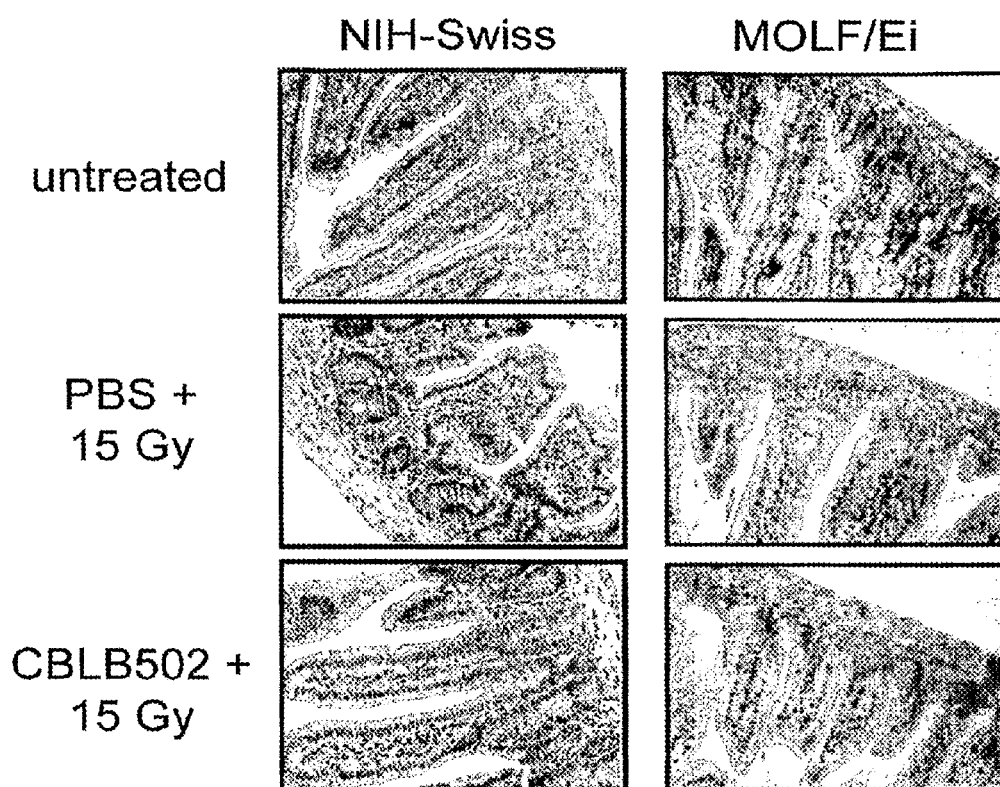


Fig. 36

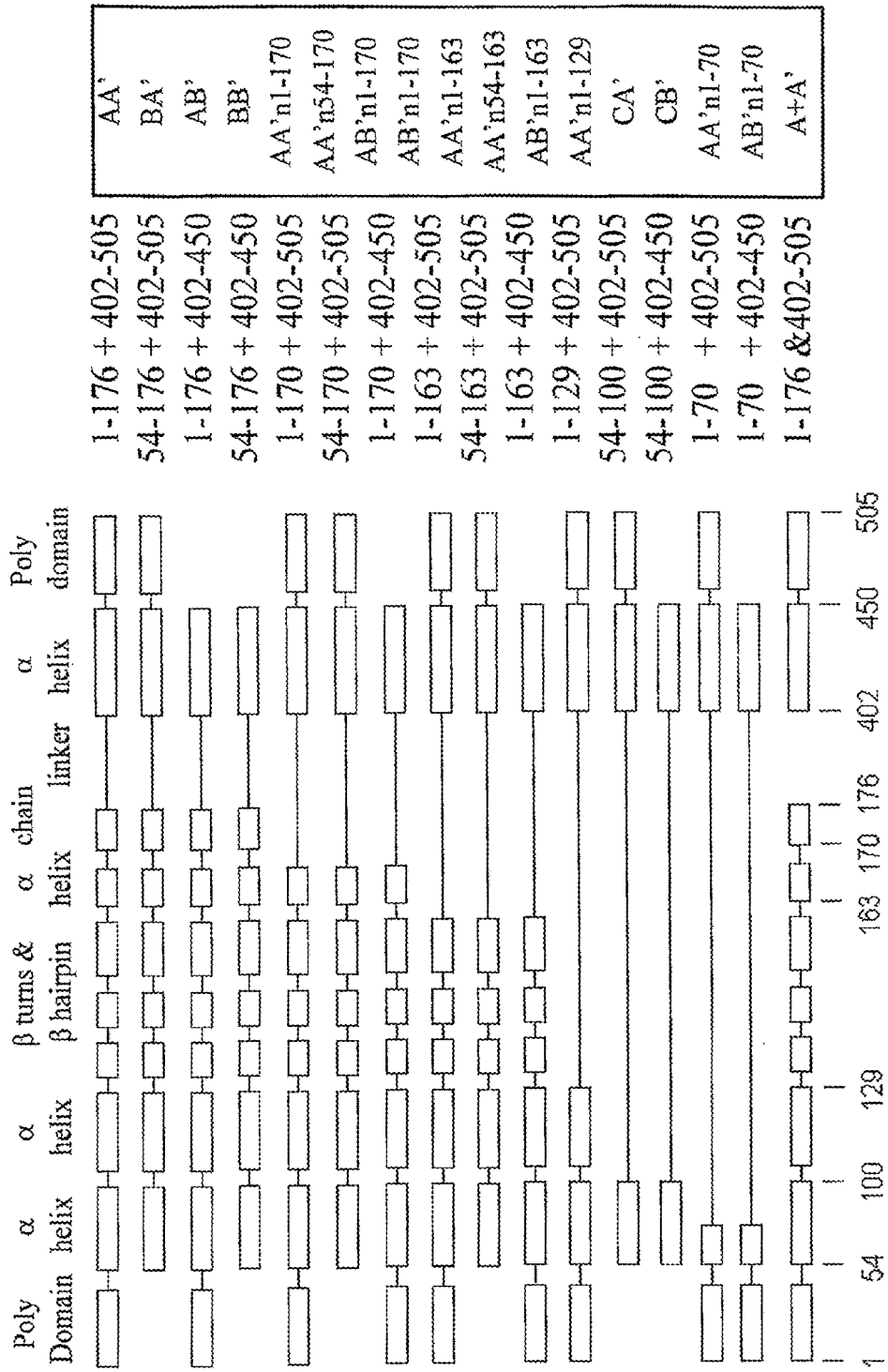


Fig. 37

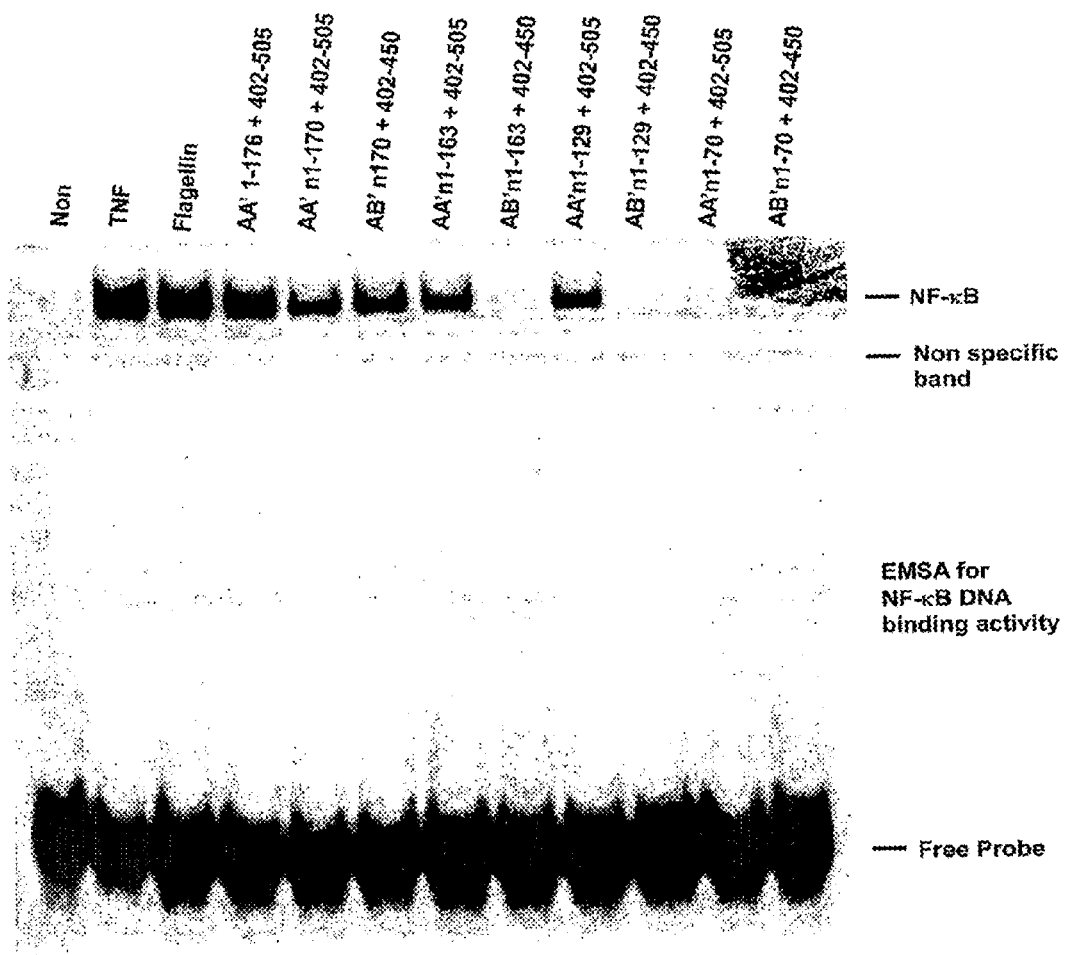


Fig. 38

AA'

Nucleotide sequence (990 bp):

```

ATGCCGGGTTCTCATCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTTACGACGATGACGATAAGGATCCGATGGCACAGTCATTAACTACA
AACAGGCTGTCGCTGTTACCCAGAATAACCTGAACAAATCTCAGTCCTCACTGAGTCC
GCTATTGAGCGTCTGCTCTGCTGCTGATCAACAGCGCGAGAGAGATGAGATGAGCAGGC
CAGGCAGTCTGTAACCGCTTCACTCTAAATATCAAAAGGCTGACTCAGGCTTCCCCTAAC
GCTAACGACGGCATTCTTATTCGCGAGACCACTGAAGGTGCGTGAAATGAAAATCAACAC
AACCTGAGCGTGTGCTGTTGAGTGTGTTCTGTCAGGCCACTAACGGGACTAACTCTGATTC
GATCTGAATATCTATCCAGATGAAATTCAACCAACGTCTGGAGAAATCAGATGCCCTTCT
AACTGAGACTCAATTAAACGGTGTAAAGTCCTCTCTCAGGCAACCGAGATGAAATCTCAG
GTTGGTCTAACGATGGTAAACATTACCATCGATCTGCAAAAAATTGAAATGAAATGAAAGC
CTGGGCTTCTGGGCTTCTGGTAAATTCCGGGAAATTCCGGTGGTGGGAAATT
CTAGACTCCATGGGTACATTAATCAATGAAACGGCTGCCGAGCCAAAGAAAAGTACCGCT
AAACCACTGGCTTCAATTGATTCTGCAATTGCAAAAGTGAACGGAGTCTGTTCTCTCTG
GGGGCAATTCAAAACCGTTTGTGATTCAAGCCATTACCAACCTTGGCAATACGGTAACCAAT
CTGAACCTGGCGCTAGCGTATCGAAGCTGACTATGCAACGGAAAGTTCTAATATG
TCTAAAGCCAGSATCTGCAAGGGCTGGTACTTCCGGTCTGGCCAGGCTAACCGGGTT
CCGAAACCGTCTCTCTTACTGCGTTAG

```

Protein sequence (329 AA):

```

MRCGSHHHHHGMASTMGGQMQGRDLYDDDDKDPMAQINTNSLTLTQNNLNKSQSSLSSAIERLSS
GLRINSAKDDAACQAIANRFTSNIKGLTQASRNANDGISAQTTGALNEINNNLQRVRELSVQATN
GTNSDSDLKSIQDEIQQRLIEIDRVSNQTOFNGVKVLSDQNMKIQVGANDGETITIDLQKIDVKSL
GLDGFDNVNSPGISGGGGILDMSMGTLINEADAACKSTANPLASIDSALKVDAVRSSLGAIQNRFD
SAITNLGNTVTMLNSARSHEIDADYATEVSNMSKAQILQQACTSVLAQANQPQNVLSLIR

```

AB'

Nucleotide sequence (825 bp):

```

ATGCCGGGTTCTCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTTACGACGATGACGATAAGGATCCGATGGCACAGTCATTAACTACA
AACAGGCTGTCGCTGTTACCCAGAATAACCTGAACAAATCTCAGTCCTCACTGAGTCC
GCTATTGAGCGTCTGCTCTGCTGCTGCTGATCAACAGCGCGAAAGACCAATGCGGGAGGC
CAGGCAGTCTGTAACCGCTTCACTCTAAATATCAAAAGGCTGACTCAGGCTTCCCCTAAC
GCTAACGACGGCATTCTTATTCGCGAGACCACTGAAGGTGCGCTGAATGAAAATCAACAC
AACCTGCAACGGTCTGGCTGACTTGTCTGTTACGGCAACTAACGGGACTAACTCTGATTC
GATCTGAAATCTATCCAGGTGAATTCAGCAACGTCGGAAAGGAAATGCAAGCGGTTCT
AACTGAGACTCAATTAAACCGCTTAAACTCTCTCTCAGGACAAACCCACATGAAATCTG
GGGGCAATTCAAAACCGTTTGTGATTCAAGCCATTACCAACCTTGGCAATACGGTAACCAAT
CTAGACTCCATGGGTACATTAATCAATGAAAGACGGCTGCCGAGGCCAGAAAAGTACCGCT
AAACCACTGGCTTCAATTGATTCTGCAATTGCAAAAGTGGACGCACTTGGTCTCTCTG
GGGGCAATTCAAAACCGTTTGTGATTCAAGCCATTACCAACCTTGGCAATACGGTAACCAAT

```

Protein sequence (274 AA):

```

MRCGSHHHHHGMASTMGGQMQGRDLYDDDDKDPMAQINTNSLTLTQNNLNKSQSSLSSAIERLSS
GLRINSAKDDAACQAIANRFTSNIKGLTQASRNANDGISAQTTGALNEINNNLQRVRELSVQATN
GTNSDSDLKSIQDEIQQRLIEIDRVSNQTOFNGVKVLSDQNMKIQVGANDGETITIDLQKIDVKSL
GLDGFDNVNSPGISGGGGILDMSMGTLINEADAACKSTANPLASIDSALKVDAVRSSLGAIQNRFD
SAITNL

```

Fig. 38 - Continued

BA'

Nucleotide sequence (831 bp):

```

ATGCGGGGTCTCATCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTACCGACGATGACGATAAGGATCCCTCACCTCTAATATCAAAGGC
CTGACTCAGGCTTCCCGTAACGCTAACGACGGCATTTCTATTGCGCAGACCACGTGAAGGT
SCGCTGAATGAAATCAACAACACCTGCAGCGTGTGCCTGAGTTGTCTGTTCAAGGCCACT
AACGGGACTTAACCTCTGATTCCGATCTGAAATCTATCTCAGGATGAAATTCAGCAACGGTCTG
GAAGAAATTCGATCGCGTCTCATGACTCAATTAAACGGTGTAAAGTCTCTCTCTCAG
GACAAACCGAGTGAATTCAGGTTGGTGTAAACGATGGTGAACCATTACCATCGATCTG
CAAAAATTCGATCTGAAAAGCCTTGGCCTTGATGGCTCAATGTTAATTCCCCGGGAATT
TCCGGTGGTGTGGTGGAAATTCTAGACTCCATGGGTACATTAACTCAATGAAGACGCGTGC
GCAGCCAAGAAAAGTACCGCTAACCCACTGGCTTCATTTGATCTGCATTGTCAAAAGTG
GACGCACTTCTCTCTGTTGGGGCAATTCAAACCGTTTGATTCAGCCATTACCAAC
CTTGGCAATACGGTAACCAATTCTGAACTCUCGGCTAGCGTATCGAAGAGATGCTCACTAT
GCAACCGAAGTTCTAAATATGCTCTAAGCGCAGATTCTGCAGCAGGCTGGTACTTCGTT
CTGGCGCAGGCTAACCAAGGTTCCGAAAACGTCCTCTTACTGCGTTAG

```

Protein sequence (276 AA):

```

MRGSHHHHHGMASMTGGQOMGRDLYDDDDKDPFTSNIKLTOASRNANDGISIAQTTEGAI
NEINNNLQRVRELSVQATNGTNSDLKSIQDEIQQRLLEIDRVSNOTQFNGVKVLSDNQMKIQVG
ANDGETITIDLQKIDVKSLGLDGFNVNSPGISGGGGIILDSMGTLINEADAACKSTANPLASIDS
ALSKVDAVRSSLGAIQNRFDASITNLGNTVTNLNSARSRIEDADYATEVSNMSKAQILOQAGTSVLA
QANQVPQNVLSSLR

```

BB'

Nucleotide sequence (666 bp):

```

ATGCGGGGTCTCATCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTACCGACGATGACGATAAGGATCCCTCACCTCTAATATCAAAGGC
CTGACTCAGGCTTCCCGTAACGCTAACGACGGCATTTCTATTGCGCAGACCACGTGAAGGT
GGCGCTGAATGAAATCAACAACACCTGCAGCGTGTGCCTGAGTTGTCTGTTCAAGGCCACT
AACGGGACTTAACCTCTGATTCCGATCTGAAATCTATCTCAGGATGAAATTCAGCAACGTC
GAAGAAATCGATCGCTTCTATCATGACTCAATTAAACGGTGTAAAGTCTCTCTCAG
GACAACCCAGATGAAATCCAGGTTGGTGTGCTAACGATGCTGAAACCATTACCATCGATCTG
CAAAAATTCGATCTGAAAAGCCTTGGCCTTGATGGCTCAATGTTAATTCCCCGGGAATT
TCCGGTGGTGTGGTGGAAATTCTAGACTCCATGGCTACATTCAATGAAAGACGCTGCC
GCAGCCAAGAAAAGTACCGCTAACCCACTGGCTTCATTTGATCTGCATTGTCAAAAGTG
GACGCACTTCTCTCTGGGGCAATTCAAACCGTTTGATTCAGCCATTACCAAC
CTTTAG

```

Protein sequence (221 AA):

```

MRGSHHHHHGMASMTGGQOMGRDLYDDDDKDPFTSNIKLTOASRNANDGISIAQTTEGAI
NEINNNLQRVRELSVQATNGTNSDLKSIQDEIQQRLLEIDRVSNOTQFNGVKVLSDNQMKIQVG
ANDGETITIDLQKIDVKSLGLDGFNVNSPGISGGGGIILDSMGTLINEADAACKSTANPLASIDS
ALSKVDAVRSSLGAIQNRFDASITNL

```

CA'

Nucleotide sequence (603 bp):

```

ATGCGGGGTCTCATCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTACCGACGATGACGATAAGGATCCCTCACCTCTAATATCAAAGGC
CTGACTCAGGCTTCCCGTAACGCTAACGACGGCATTTCTATTGCGCAGACCACGTGAAGGT

```

Fig. 38 - Continued

GGCGCTGAATCAAACAAACACTGCAGCGTGTGCGTGAGTTGTCTGTTCAAGGCCACT
TCCCCGGGAATTCCGGTGGTGGGAAATTCTAGACTCCATGGGTACATTATCAAT
GAAGACCGCTGCCGCAGCCAAGAAAAGTACCGCTAACCCACTGGCTTCATTGAAATTCTGCA
TTGTCAAAAGTGGACGCAGITCGTTCTCTCIGGGGGCAATTCAAAACCGTTTGATTCA
GCCATTACCAACCTTGGCAATACGGTAACCCATCTAACACTCCGCGCTAACCGTATCGAA
GATGCTGACTATGCAACGGAAGTTCTAATATGTCATAACCGCAGATTCTGAGCAGGCT
GGTACCTCGTTCTGGCGCAGGCTAACCAAGGTTCCGCAAACGTCCTCTTTACTGCGT
TAG

Protein sequence (200 AA):

MRGSHHHHHHGMASMTGGQQMGRDLYDDDDKDPFTSNIKGLTQASRNANDGISAQTTEGALNEINN
NLQRVRELSPVQATSPGISMGGGGILDSMGTLINEADAAAKSTANPLASIDSALKVDAVRSSLGAI
QNRFDSAINTNLGNTVTNLNSARSRIEDADYATEVSNMSKAQILOQAGTSVLAQANQPQNVLSLLR

CB'

Nucleotide sequence (438 bp):

ATGCCGGGTTCTCATCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTACGACGATGACGATAAGGATCCGTTACTTCTATATCAAAAGGC
CTGACTCAAGCTTCCGTAACCTAACGAAAGGCAATTCTATTCGCGCAGACCAACTGAAGGT
GCGCTGAATCAAACACACCTCCAGCGTGTCCGTGAGTTGTCTTCAAGGCCACT
TCCCCGGGAATTCCGGTGGTGGGAAATTCTAGACTCCATGGGTACATTCAAT
GAAGACCGCTGCCGCAGCCAAGAAAAGTACCGCTAACCCACTGGCTTCATTGATTCTGCA
TTGTCAAAAGTGGACGCAGITCGTTCTCTCIGGGGGCAATTCAAAACCGTTTGATTCA
GCCATTACCAACCTTTAG

Protein sequence (145 AA):

MRGSHHHHHHGMASMTGGQQMGRDLYDDDDKDPFTSNIKGLTQASRNANDGISAQTTEGALNEINN
NLQRVRELSPVQATSPGISMGGGGILDSMGTLINEADAAAKSTANPLASIDSALKVDAVRSSLGAI
QNRFDSAINTNL

A

Nucleotide sequence (639 bp):

ATGCCGGGTTCTCATCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTACGACGATGACGATAAGGATCCGATGACAAAGTCATTAAATACA
AACAGCTCTGCGTGTGACGCCAGAAATAACCTGAAACAAATCTCAGTCCCTACTGAGTTCC
SCTATTGAGCTCTGTCCTGCGTGTGCGTATCAACBGCUGCGAAAGACGATGCCGAGGC
CAGCGGATTCATAACCGCTTCRACTCTATAATCAAAAGGTTCTGACTCAGGCTCCCGTAAAC
GCTAACGACGGCATTCTATTCGCGCAGACCACTGAAGGTGGCTGAATGAAATCAACAAAC
AACCTGCAAGCTGTGCGTGTGACTCTGCTGCTGCGCCACTAACGGGACTTAACCTCTGAATTC
GATCTGAAAATCTATCCAGGATGAAATTCAAGAACGCTTGAGAGAAAATGATCGCGTTCT
AATCAGACTCAATTAAACGGTGTAAAGTCTGCTCAGGACAAACCGAGATGAAAATCCAG
GTTGGTGTAAACGATGGTGAACCAATTACCAUCGATCTGCCAAAAATTGATGTGAAAAGC
CTTGGCCCTGATGGGTTCAATGTTAATTCCCCGGGATGA

Protein sequence (212 AA), last three amino acids are derived from primer and pRSETb polylinker:

MRGSHHHHHHGMASMTGGQQMGRDLYDDDDKDPMAQVINTNSLTLTQNNLNKSQSSLSSAIERLSS
GLRINSAKDAACQATANRFTSNIKGLTQASRNANDGISAQTTEGALNEINNNLQRVRELSVQATN
GTNSDSDLKSIQDEIQQRLEELDRVSNQTOFNGVKVLSDQNQMKIQVGANDGETITIDLQKIDVKSL
GLDGFNVNSFG

Fig. 38 - Continued

B

Nucleotide sequence (480 bp):

```

ATGCCGGGTTCTCATCATCATCATCATCATCGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGGATCTGTACGACGATGACGATAAGGAACTCGTTCACTTCTAATATCAAAGGT
CTGACTCAGGCTTCCGTAACGCTAACGACGUCATTCTTATTCGCGCAGACCAACTGAAGGT
GCCTGAATGAATGAAACAAACCTCCACCGTGTGGCTGAGTTGTCAGGCCACT
AACGGGACTAACCTGATTCCGATCTGAATACTATCCAGGATGAAATCAAGAACGCTG
GAAGAAATCGATGGCTTCTAATCAGACTCAATTAAACGGTGTAAAGTCTGTCAG
GACAAACCAGATGAAATCCAGGTGGCTAACGATGGTAACCCATTACCATCGACCTG
CAAAAAATTGATGTGAAAAGCCTTGCTATGGCTTCAATGTTAATTCAGGATG

```

Protein sequence (159 AA), last three amino acids are derived from primer and pRSETb polylinker:

```

MRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPFTSNIKLTOASRNANDGISIAQTTEGALNEINN
NLQRVRELSVQATNGTNSDILKSIQDEIQQRLEEDRVSNQTFNGVKVLSDQNQMKIQVGANDGE
TITIDLQKIDVKSLGLDFNVNSPG

```

C

Nucleotide sequence (252 bp):

```

ATGCCGGGTTCTCATCATCATCATCGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGGATCTGTACGACGATGACGATAAGGATCCGTTCACTTCTAATATCAAAGGT
CTGACTCAGGTTCCGTAACGCTAACGACGGCATTTCTATTCCGCGACACCACTGAAGGT
GCCTGAATGAATGAAACAAACACCTGCAAGCGTGTGCGTGAGTTGTCAGGCCACT
TCCCCGGGATGA

```

Protein sequence (83 AA), last three amino acids are derived from primer and pRSETb polylinker:

```

MRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPFTSNIKLTOASRNANDGISIAQTTEGALNEINN
NLQRVRELSVQATSPG

```

GST-A'

Nucleotide sequence (1038 bp), GST highlighted:

```

ATGTCCTCTATACTAGGTTATTGAAAATTAAAGGGCTTGTGCAACCACTCGACTCTT
TTGGAAATATCTGAAAGAAAATATGAAAGAGCATTGATGAGGCCGATGAAGGTTGATAAAA
TGGCGAAACAAATTTGAAATTGGCTTCCGAGTTCCCAATCTCCCTTATATATGAT
GGTGAATGTTAAATTAAACACAGCTATGGCCATCATACTTATATAGCTGACAAGCACAAAC
ATGTCGGGTGGTTCTCAAAAGACGCTGCAAGAGATTCTCATGCTTGAAGGAGCGTTTGTG
GATATTAGATAACGGTGTTCGAGAAATTGCAATAGTAAAGACTTGTAAACTCTCAAAAGTT
GATTTCTCTAGCAAGCTACCGTAAATGCTGAAATGTTCCAAGATGCTTAAATGTCATAAA
ACATAATTAAATGGTGATCATGTAACCCATCCTGACTCTCATGTTATGACGCTCTGAT
GTTGTTTATACGGACCCATGGCTGGATGGCTTCCAAATTAGTTAGTTTGTAAA
AAACCTATTGAAAGCTATCCCACAAATTGATAAGTACTTGAATCCAGCAAGTATATAGCA
TGGCTTTTGCAGGGCTGGCAAGCCACGTTGGTGGTGGCGACCATCCTCCAAATCTGGAT
CTGGTTCCGGCTGGATCCCCGGAAATTCCGGTGGTGGTGGTGGAAATCTAGACTCCATG
GGTACATTAACTAAAGAACGCTGCCCGCAGCCAAGAAAAGTACCCCTAACCCACTGGCT
TCAATTGATTCTGCATTGCAAAAGTGGACGCAAGTTGGCTTCTTGTGGGGCAATTCAA
AACCGTTTGTGATTCAGCCATTACCAACCTGGCAATACGGTAACCAATCTGAACCTGGCG
CGTAGCCGATATGGAAAGATGCTGACTTATGCAACGGAACTTCTAATATGCTAAAGGGCAG
ATTCTGCAGCAGGTGGTACTTCGGTGGCTGGCTAACCGAGTTCCGCAAAACGTC
CTCTTTTACTGCGTTAG

```

Protein sequence (345 AA):

Fig. 38 - Continued

MSPILGYWKIKGLVQPTTRLLLEYLEEKYEEHLYERDEGDKWRNKKFELGLEFPNLPYYIDGDVKLTQ
 SMAIRYIADKHNMLGGCPKERAETISMLEGAVLDIRYGVSRAYSKDFETLKVDFLSKLPEMLKMFE
 DRLCHKTYLNCDHVTHPDFMLYDALDVVLYMDPMCLDAFPKLVCFKKRIFALEQIDKYLKS8KYIAW
 PLQGWQATFGGGDHPPKSDLVPRGSPGISGGGGILDMSGTLINEDAAAACKSTANPLASIDSALK
 VDAVRSSLGAIQNRFDSAITNLGNNTVNLSARSRIEDADYATEVSNMSKAQILQOAGTSVLAQANO
 VPQNVLSLLR

GST-B'

Nucleotide sequence (873 bp), GST highlighted:

ATGTCQQCTTACTAGGTTATTGGAAAATAAGGCCCTGTGCAACCCACTCGACTCTT
TTGAAATATCTTGAAAGAAAATATGAAAGAGCATTGTATGACGCCATGAAGGTGAA
ATGGCAACAAAAGCTTAAATGCGTTGGAGTTTCCCAATUTYKCTTAAATATCTG
GGTGATGTTAAATTAACACAGCTATGGCATCATACGTTATACTGACAAGCACAC
ATGGGGCTGTGTGCAAAGAGCGTGCAGAGATTTCAGCTTGAGAGGGTTTG
GATATTAGATAACGGTGTTCAGAATTCGATATAGTAAAGACTTGAAACTCTCAAGCT
CATTTCTTACGACCTACCTGAAATGCTGAAATGCTGAAATGCTGATGCTTATGCTAA
ACATTTAAATGGTCAATCATGAAACCCATCTGACTTCACTGTTGACGCTCUTGAT
GTTTTTAAATCATGGACCCAATGCTGGATGCGTTCCAAATATTACCTTGTGTTAA
AACGTTATGAAAGCTATCCCACRAATTGATAAGTACTTGAATCCAGCAAGTATATAGCA
TGGCTTTGCAAGGGCTGGCAAGCACCGTTGGGGGGCGSACCACCTCCAAATCGGAT
CTGGTTCCGGGGATCCCCGGAAATTCCGGTGGTGGTGGATCTAGACTCCATG
GGTACATTATCATGAAAGACCGTGCACGCAAGAAAGTACCGCTAACCCACTGGCT
TCAATTGATTCGGATTGTCAAAGTGGACGCACTTCGTTCTCTGGGGCAATTCAA
AACCGTTTGATTGAGCCATTACCAACCTTAG

Protein sequence (290 AA):

MSPILGYWKIKGLVQPTTRLLLEYLEEKYEEHLYERDEGDKWRNKKFELGLEFPNLPYYIDGDVKLTQ
 SMAIRYIADKHNMLGGCPKERAETISMLEGAVLDIRYGVSRAYSKDFETLKVDFLSKLPEMLKMFE
 DRLCHKTYLNCDHVTHPDFMLYDALDVVLYMDPMCLDAFPKLVCFKKRIFALEQIDKYLKS8KYIAW
 PLQGWQATFGGGDHPPKSDLVPRGSPGISGGGGILDMSGTLINEDAAAACKSTANPLASIDSALK
 VDAVRSSLGAIQNRFDSAITNL

AA'n1-170

Nucleotide sequence (972 bp):

ATGCGGGGTTCTCATCATCATCATGGTATGGCTAGCATGACTGGGGACAGCAA
ATGGGTCCGGATCTGTAACGACGATGACGATAAGGATCCGATGGCAACAGCTATTAA
AAACAGCCTGTCCTGTTGACCCAGATTAACCTAAACAAATCTCAGTCTCTACTGAGGTTCC
GCTATTGAGGCTCTGCTCTGCTGCTGCTGCTGATGCGTATCAGCAGGCSAAAGACGATGGCGAGGC
CAGGGCATTGGCTACOGCTTCACTCTTAATATCAAAAGGCCGACTCAAGCTTCCCGTAAC
GCTAACGACGCCATTCTATTGGCGAAGACACTGAAAGTGCCTGGAATGAAATCAAAAC
AACTGCAAGGGTGTGGGTGAGTTGCTGTCTGAGGCCACTAACGGGACTTAACCTCTGATTCC
GATCTGAAATCTATCCAGGATGAAATTTCAGGAAAGTCTGGAAAGAAATCGATGCGGTTCT
AAATCAGACTCAATTAAACGGGTAAAGTCTCTCTCAGGACAAACCGGATGAAAATCCAG
CTTGGCTTAACGATGCTGAAACCATTAACCATGAAATGCAAAATGATGAAAGC
CTTGGCCTTAATCCGGGAATTCCGGTGGTGGGAATTCTAGACTCCATGGCTACA
TTAATCTAATGAAAGACGCTGCCGAGCCAAGAAAAGTACCGCTAACCCACIIGGCTTCATT
GATCTGCAATGTCATAAGTGGACCGTGTGTTCTGCTGSSGGCAATTCAAAACCGT
TTTGATTCAGCCATTACCAACCTTGCAATACGGTAACCAATCTGAACTCCGGCGTAGC
CGTATGCAAGATGCTGACTATGCAACGGAGTTCTTAATATGTCATAAGCGCAGATTCTG
CAGCAGGGCTGGTACTTCGGTTCTGGGGCAGGGCTAACCGAGGTTCCGCAABACGTCTCTCT

Fig. 38 - Continued

TAACTGGCGTTAG

Protein sequence (323 AA):

MRCGSHHHHHHGMASMTGGQQMGRDLYYDDDDKDPMAQVINTNSLSLLTQNNLNKSQSSLSSAIERLSS
GLRINSAKDDAAGQAIANRFTSNIKGLTQASRNANDGISAOTTEGALNEINNNIQRVRELSVQATN
GTNSDSDLKSIQDEIQORLEEIDRVSNQTFNGVKVLSDQNQMKTQVGANDGETITIDLQKIDVKS
GLIPGIGSGGGGILDMSGTLINEDAAAACKSTANPLASIDSALKVDAVRSSLGAIQNRFDSAITNL
GNTVTNLNSARSRIEDADYATEVSNMSKAQILQOAGTSVLAQANQVPQNVLSSLR

AA'n1-163

Nucleotide sequence (951 bp):

AATCGGGGGTTCTCATCATCATCATCATGGTATGGCTAGCATGACTGGTAGCACAGCAA
ATGGGTCTGGATCTGTACCGACGATGACGATAAGGAATCCGATGGCACAAAGTCATTAATACA
AAACAGCCTGCGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTG
GCTATTGAGCGTCATGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTGCTG
CAGCGGATTCGCTAACCGCTTCACTTCTAATATCAAAGGCCGACTCAGGGCTTCCCCGTAAC
GCTAACCGACGGCAATTCTATCCGGAGACCACTGAAAGGTGGCTGAAATGAAATCAACAAAC
AAACCTGCAAGCGTGTGCCTGAGGTTGTCAGGCCACTAACCGGACTAACTCTGATTTCC
GATCTGCAAAATCTCAGGATGAAATTCAGCAACGCTCTGGAAAGAAATCGATGGCTGTTCT
AAATCAGACTCAATTAAACGGTGTAAACTCTCTCTCAGGACAACCAACGAGATGAAATATCCAG
GTGGTGCTAACGATGGTAAACCATTAACCATCGATCTGCAAAAAATTATCCGGGAATT
TCCGGTGGTGGTGGTGGAAATCTAGACTCATGGCTACATTAAATCAATGAAAGACGCTGCC
GCAGCCAAGAAAATCTACCGCTAACCCACTGGCTAACATGATCTGCTATGCTAAAGTG
GACGGCAGTTCTCTCTCTCTGGGGGCAATTCAAACCGGTTGATTCAAGGCAATTACCAAC
CTTGGCAATTACGGTAACCAACTCTGAACTCCGGCGCTAGCCGATGCAAGCTGCTGACTAT
GCACACGGAACTTCTAAATATGCTCTAAAGCGCAAGATTCTGCAAGCAGGCTGGTACTTCGTT
CTGGCGCAGGCTAACCGGTCTCGCAAAACGTCCTCTCTTACTGCGTTAG

Protein sequence (316 AA):

MRCGSHHHHHHGMASMTGGQQMGRDLYYDDDDKDPMAQVINTNSLSLLTQNNLNKSQSSLSSAIERLSS
GLRINSAKDDAAGQAIANRFTSNIKGLTQASRNANDGISAOTTEGALNEINNNIQRVRELSVQATN
GTNSDSDLKSIQDEIQORLEEIDRVSNQTFNGVKVLSDQNQMKTQVGANDGETITIDLQKIDVKS
GGGGGILDMSGTLINEDAAAACKSTANPLASIDSALKVDAVRSSLGAIQNRFDSAITNLGNTVTNL
NSARSRIEDADYATEVSNMSKAQILQOAGTSVLAQANQVPQNVLSSLR

AA'n54-170

Nucleotide sequence (813 bp):

ATGCGGGGGTTCTCATCATCATCATCATGGTATGGCTAGCATGACTGGTAGCACAGCAA
ATGGGTCTGGATCTGTACCGACGATGACGATAAGGAATCCGTTACITCTAATATCAAAGC
CTGACTCAGGCTTCCGTAACGCTAACGACGGCAATTCTATTGCGCAGAACACTGAAAGGT
GGCGCTGAATGAAATCAACAAACCTGCAAGCGCTGCTGCGTGAATTGTCAGGGCACT
ACCGGACTRACTCTGATTCGGATCTGAAATCTCAATCCAGGATGAAATTCAAGAACGCTG
GAAGAAATGATGGTTCTCTAATCAGACTCAATTAAACGGTTAAAGTCCTCTCTCAG
GACAACCAGATGAAAATCAGGTTGGTGTAAACGATGGTGAACCAATTACCATCGATCTG
CAAAAKATGATGATGAAAACCTTGGCTTATCCGGGAAATTCCGGGGTGGTGGTGG
ATTCCTAGACTCTCATGGTATCATTTAATGAAAGACCTCCCGCAGCCAAGAAAAGTACC
CTTAAACCCACTGGCTTCATGATGATGATGATGATGATGATGATGATGATGATGATG
CTGGGGGCAATTCAAACCGGTTGATCTGCAATTACCAAACCTTGGCAATTACGGTAAAC
ATCTGAAACTCCGGCGTGGCCGATGAGAGATCTGACTATGCAACCGGAAGTTCTAAT
ATGTCATAAAGCGCAGATTCTGCAAGCAGGCTGGTACTTCGTTCTGGCGAGGCTAACCG
CTTCGCAAAACGTCCTCTTACTGCGTTAG

Fig. 38 - Continued

Protein sequence (270 AA):

MRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPPFTSNIKGLTOASRNANDGISIAQTTEGALNEINN
NLQRVRELNVQATNGTNSDSLKSIDQDEIQQRLEEDRVSNQTOFNGVKVLSDNQMKIQVGANDGE
TITIDLQKIDVKSLGLIPGISGGGGILDSMGTLINEDAAAKSTANPLASIDSALKVDAVRSSL
GAIQNRFDSAITNLGNTVTNLNSARSRIEDADYATEVSNMSKAQILOQAGTSVLAQANQVPQNVLSL
LR

AA'n54-163

Nucleotide sequence (792 bp):

ATGCCGGGGTTCTCATCATCATCATGGTATGGCTAGCATCACTGGTGACAGCAA
ATGGGTGGGATCTCTGACAGCAGATGGATTCAGGATCCGGTCACCTCTAATATCAAGGC
CTGACTCAGGCTTCCCGTAACGCTAACGACGGCATTTCTATTCGGCAGACCACTGAAGGT
SGCGCTGAATGAATCAACAACAAACCTGCAGCGTGTGGTGGAGTTGCTGTTCTCAGGGCACT
AACCGGACTAACCTGATTCCGATCTGAAATCTATCCAGGATGAAATTCAAGCAACGCTTG
GAAGAAATCTGATCCGGCTTCTCTACAGACTCAATTAAACGGTGTAAAGTCCTCTCAG
GACAAACGAGATGAAAATCCAGGTTGGTGTAAACGATGGTGAACACATTACCATGATCTG
CAAAATAATTATCCGGGAATTCCGGTGGTGTGGTGGAAATCTAGACTCCATGGGTACA
TTAATCAATGAAGACGGTGGCGCAAGAAAAGTACCGCTAACCCACTGCTTCATCAATT
GATCTGCGATGCAAAAGTGAGCAGTTCTCTGGGGCAATTCAAACCGT
TTTAAATTAGCTCATTAACCAACCTTGGCAATACGTTAACCAATCTGAACCTGGCGCTTAC
CGTATCCAAGATCTGACTATGCAACCGAAAGTTCTAATATCTCTAAAGGCCAGATCTG
CAGCAGGCTGGTACTTCCGTTCTGGCGCAGGCTAACAGGTTCGGCAAAACGTCCTCTCT
TTACTGCGTTAG

Protein sequence (263 AA):

MRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPPFTSNIKGLTOASRNANDGISIAQTTEGALNEINN
NLQRVRELNVQATNGTNSDSLKSIDQDEIQQRLEEDRVSNQTOFNGVKVLSDNQMKIQVGANDGE
TITIDLQKIDVKSLGLIPGISGGGGILDSMGTLINEDAAAKSTANPLASIDSALKVDAVRSSL
GAIQNRFDSAITNLGNTVTNLNSARSRIEDADYATEVSNMSKAQILOQAGTSVLAQANQVPQNVLSLLR

AB'n1-170 (or AA'n1-170c402-450)

Nucleotide sequence (807 bp):

ATGCCGGGGTTCTCATCATCATCATGGTATGGCTAGCATCACTGGTGACAGCAA
ATGGGTGGGATCTCTGACAGCAGATGGATTCAGGATCCGGTCACAGCTATTAATACA
AAACAGCTGCGCTTGTGACGGCAAAATTAACCTAACAAATCTCACTGCTCACTGAGTTCC
GCTATTGACGCTCTGCTCTGGCTGCTGATCACAGCGCGAAAGACGATGCGCAGCC
CAGGGGATCTAACCGCTTCACTCTAATATCAAAAGGCTGACTCAGGCCTCCGTAAC
GCTAACGCGACGGCATTTCTATTCGGCAGACCACTGAAAGGTGCCTGATGAAATCAACAC
AACTGCGACGGCTGCGTGAAGTTCTCTGAGGCGACTAACGGGACTAACCTGATTTCC
GATCTGAAATCTATTCAGGATGAAATTCAAGCAACGCTCTGAAAGATCGATCCGTTCT
AATCAGACTCTATTAAACGGTGTAAAGTCTCTCTCAGGACAACCGAGATGAAATCCAG
GTTCGGCTTATCCGGGAATTTCGGTGTGGAAATTCTAGACTCCATGGGTACA
TTAATCAATGAAGACCCCTGCCCAACCGAAAGAAAAGTACCGCTAACCCACTGCGCTCAATT
GATCTGCGATTGTCAGGCCATTACCAACCTTGTAG

Protein sequence (268 AA):

MRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPPFTSNIKGLTOASRNANDGISIAQTTEGALNEINNNLQRVRELNVQATN
GLRINSAKDDAACQAIANRFTSNIKGLTOASRNANDGISIAQTTEGALNEINNNLQRVRELNVQATN

Fig. 38 - Continued

GTNSDSLKSIQDEIQRLEEIDRVSNQTFNGVKVLSDNQMKIQVGANDGETITIDLOKIDVKSL
GLIPGISGGGGGILDMSGTLINEDAAAACKSTANPLASIDSALKVDAVRSSLGATQNRFDSAINTNL

AB'n1-163 (or AA'n1-163c402-450)

Nucleotide sequence (786 bp):

```
ATCGGGGGTTCTCATCATCATCATCATGGTATGGCTAGGCATGACTGGTGGACAGCAA
ATGGGTCTGGGATCTGTACCGACGATGACGATAAGGATCCGATGGCACAGTCATTAAATACA
AACAGCCCTGTGCTGTTGACOCAGAATAACCTGAACAAATCTCAGCCTCACTGAGTTCC
GCTATTGAGCGCTGTGCTCTGGTCTGGTATCAACAGCGGAAAGACCGAAGGGCAGGC
CAGGGGATTGCTAACCGCTTCACTTCTTAATATCAAAAGGCTGACTCAGGCTTCCGTAAC
GOTAAACGACGCCATTTCTATGGCAGACCACTGAAAGGTGCGCTGAATGAAATCAACACAC
AACCTGAGCGCTGTGCTGAGTGTGCTGAGGUCACTAAACCTGAGTTCC
GATCTGAAATCTTCCAGATGAAATCTCAGCAGCAGCTGGAAGAAATCCGATCCGTTCT
AAATCAGACTCAATTAAACGTTAAACCTCTCTCAGGACAAACAGATGAAAATCCAG
GTGGTGTAAACGATGGTAAACCATCGATCTGCAAAATATCCGGAAATT
TCCGGTGTGGTGGAAATCTGACTCTCATGGTACATTAAATCAATGAAGACGCTGCC
GAGGCCAAGAAAAGTACCGCTAACCCACTGGCTTGATTGATTCTGCAATTGTCAAAAGTG
GACCGAGTTCTCTCTCTGGGGCAATTCAAAACGTTTGATTCAAGCCATTACCAAC
CTTTAG
```

Protein sequence (261 AA):

```
MRGSHHHHHHGMASMTGGQOMGRDLYYDDDDKDPMAQVINTNSLSSLTONNLNKSOSSLSSAIERLSS
GLRINSAKDDAACQAIAANRFTSNIKGLTOASRNANDGISAQTTEGALNEINNNLQRVRELSVQATN
GTNSDSLKSIQDEIQRLEEIDRVSNQTFNGVKVLSDNQMKIQVGANDGETITIDLOKIIIPGIS
GGGGGILDMSGTLINEDAAAACKSTANPLASIDSALKVDAVRSSLGATQNRFDSAINTNL
```

AA'n1-129

Nucleotide sequence (849 bp):

```
ATCGGGGGTTCTCATCATCATCATCATGGTATGGCTAGGCATGACTGGTGGACAGCAA
ATGGGTCTGGGATCTGTACCGACGATGACGATAAGGATCCGATGGCACAGTCATTAAATACA
AACAGCCCTGTGCTGTTGACOCAGAATAACCTGAACAAATCTCAGCCTCACTGAGTTCC
GCTATTGAGCGCTGTGCTCTGGTCTGGTATCAACAGCGGAAAGACGATGCGGAGGC
CAGGGGATTGCTAACCGCTTCACTTCTTAATATCAAAAGGCTGACTCAGGCTTCCGTAAC
GCTAAACGACGCCATTTCTATGGCAGACCACTGAAAGCTGCGCTGAAATGAAATCAACACAC
AACCTGAGCGCTGTGCTGAGTGTGCTGAGGCCACTAAACGCGACTAACTCTGATTCG
GATCTGAAATCTATCCAGGATGAAATTGAGCAACGTCGGCAAGAAATCGATCGCTTTCT
AAATCAGATGAAACCGGGAAATTCTCGGTGGTGGGAAATCTAGACTCTCATGGTACATTA
ATCAATGAAACCGCTTCCGGCAGGAAAGTACCGCTAACCCACIGGCTTCAATGAT
CTTGCAATTGTCATAAGTGGACGGAGTTCTCTCTCTGSGGGCCTGATCAAAACGTTT
GATTCAGCCATTACCAACCTGGCAATTACGGTAACCAATCTGAAACTCCGGCGTAGCCG
ATCGAAGATGCTGACTATGCAACGGAAAGTTCTAATATGCTAAAGCGCAGATTCTGCAAG
CAGGGCTGGTACTTCCGTTCTGCGCGCAGGCTAACAGGTTCCGCAAAACGTCCTCTCTITA
CTCGCTTGTAG
```

Protein sequence (282 AA):

```
MRGSHHHHHHGMASMTGGQOMGRDLYYDDDDKDPMAQVINTNSLSSLTONNLNKSOSSLSSAIERLSS
GLRINSAKDDAACQAIAANRFTSNIKGLTOASRNANDGISAQTTEGALNEINNNLQRVRELSVQATN
GTNSDSLKSIQDEIQRLEEIDRVSNQTFNGVKVLSDNQMKIQVGANDGETITIDLOKIIIPGIS
GGGGGILDMSGTLINEDAAAACKSTANPLASIDSALKVDAVRSSLGATQNRFDSAINTNL
ALSALKVDAVRSSLGATQNRFDSAINTNLGNVTNLNSARSRIEDADYATEVSNSMSKAQILQQAGTSVIA
QANQVPQNVLSLLR
```

Fig. 38 - Continued

AA'n54-129

Nucleotide sequence (690 bp):

```

ATGCCGGGGTTCTCATCATCATCATCATCATGGTATGGCTAGCATGGTGACAGCAA
ATGGGTCCGGGATCTGTAACGACGGATGACGATAAGGATCCGGTCACTCTAATATCAAAGGC
CTGACTCAGGCTTCGGTAAACGCTAACGACGGCATTTCATTCGGCGACACCACTGAAGGT
GCCTGAAATGAATCAACAAACCTGCAAGCGTGTGGTGAAGTTCTCTGTTCAAGGOCACT
AACGGGACTAATCTGATTCCGATCTGAATCTATCCAGGGATGAAATTCAAGAACCTCTG
GAAGAAATCGATCGCGTTCTAATCAGATCCCGGAATTCCCGTGGTGGTGGTGGAAATT
CTAGACTCCATGGGTACATTAATCAATGAAGAGACCGTGCAGCAGCCAAGAAAAGTACCGCT
AACCCACTGGCTTCATCTGATTCGATTGTCATTAACCTGGATCCAGTTCGGTCTTCCTG
GGGGCAATTCAAAACCGTTTGATTAGCCATTACCAACCTTCGCAATACGGTAACCAAT
CTGAACTCCCGCGTAGCCSTATCGAAGATGCTGACTATGCAACGGAAGTTCTAATATG
TCTAAGCGCAGAATCTGCAGCAGGCTGGTACTTCGGTCTGGCGCAGGCTAACAGGTT
CCGCAAAACGTCCTCTTACTGGCTTAG

```

Protein sequence (229 AA):

```

MRGSHHHHHGMASMTCCQQMGRDLYDDDDKDPFTSNIKGLTQASRNANDGISIAQTTEGALNEINN
NLQRVRELSQLVQATNGTNSDSDLXSLIQDEIQQRLEEDRVSQIPGIGGGGGILDMSGTLINEDAAA
AKKSTANPLASIDSALKVDAVRSSLGAIQNRFDSAINTNLGNTVTNLNSARSRIEDADYATEVSNMS
KAQILQQAGTSVLAQANQVPQNVLSLLR

```

AB'n1-129

Nucleotide sequence (684 bp):

```

ATGCCGGGGTTCTCATCATCATCATGGTATGGCTAGCATGGTGACAGCAA
ATGGGTCCGGGATCTGTAACGACGGATGACGATAAGGATCCGGTCACTCTAATATCAAACA
AACAGCCTGTCGCTTGTGACCCAGAATAACCTGAAACAAATCTGACTCCTACTGAGTTCC
GCTATGAGCCCTCTGTCCTCTGGTCTGCGTATCACAGCGCGAGAACGAAGCGGAGGC
CAGCCGATTGGCTCACTCTAATATCAAAAGGCCACTCAGGCTTCCCCGTAAC
GTAACGAGCGCAATTCTATGGCGAACACTGAGGTGGTGAATGAAATCAGAAC
AACCTGCAGCGTGTGGTCACTGCTGAGTTGTCCTTCAGGCCACTAACGGGACTAACTCTGATTC
GATCTGAAATCTATCAGGATGAAATTCAAGCAACGCTCTGAAGAAATCGATGGCGTTCT
AATCAAGATCCCGGAATTTCGCGGTTGGTGGTGAATTCTGAGACTCAGGGTACATIA
ATCAATGAAGAGCGTCCCGCAGCAGAAAAGTACGGCTAACCCACTGGCTTCATTTGAT
TCTGCATTTGCAAAACGGACGGCAGTTCTCTCTGGGGCAATCAAAACGGTTT
GATTCAGCCATTACCAACCTTAG

```

Protein sequence (227 AA):

```

MRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPMAQVINTNSLSSLTQNNLNKSQSSLSSAIERLSS
GLRINSAKDDAAGQATIANRFTSNIKGLTQASRNANDGISIAQTTEGALNEINNMLQRVRELSQLVQATN
GTNSDSDLKSIQDEIQQRLEEDRVSQIPGIGGGGGILDMSGTLINEDAAAACKSTANPLASIDS
ALKVDAVRSSLGAIQNRFDSAINTNL

```

AB'n54-129

Nucleotide sequence (525 bp):

```

ATGCCGGGCTCTCATCATCATCATGGTATGGCTAGCATGGTGACAGCAA
ATGGGTCCGGGATCTGTAACGACGGATGACGATAAGGATCCGGTCACTCTAATATCAAAGGC
CTGACTCAGGCTTCGGCTAACGCTAACGACGGCATTTCATTCGGCGACACCACTGAAGGT
GCCTGAAATGAATCAACAAACCTGCAAGCGTGTGGTGAAGTTCTGTTCAAGGCCACT

```

Fig. 38 - Continued

AACGGGACTAACCTCTGATTCCGATCTGAAATCTATCCAGGATGAAATTCAAGCAACCTCTG
GAAGAAATCGATCGCGTTCTAATCAGATCCCGGGAAATTTCGGTGGTGGGTGGAAATT
CTAGACTCCATGGGTACATTAATCANTGAAGACGCTGCCAGSCCAAGAAAAGTACCGCT
AACCCACTGCTCAATTGATCTGCATTCTCAAAGTGACCCAGITCGTCTTCTCTG
GGGCAATTCAAAACCGTTTGATTCAGCCATTACCAACCTTAG

Protein sequence (174 AA):

MRGSHHHHHGMASMTGGQQMGRDLYYDDDKDPMAQVINTNSLTLQNNLNKSQSSLSSAIERLSS
NLQRVRELSVQATNGTNSDSDLKSIODEIQRLEEDRVSNQTPGISGGGGILDMSGTLIN
AKKSTANPLASIDSALSKVDAVRSSLGATQNRFDSAITNL

AA'n1-100

Nucleotide sequence (762 bp):

ATGCCGGGTTCTCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTGACGGATGACGATAAGGAATCCGATGGCACAGTCATTAAACA
AACAGCCCTCGCTGGGATCTGTGACGGATGACGATAACCTGAACAAATCTCAGTCCTACTGAGTTCC
GCTATTGAGGCTCTGCTCTGGTCTGGTCTGAGGCTGACTCTGAGGCTTCCGGTAAAC
GCTAACGACGGCATTCTTATGGCGCAGACCACTGAAGGGTGGCTGAANGAATCAACAC
AACCTGCAACGGCTGGTGTGAGGTTGCTGAGGCTACTATCCGGGAATTCCGGTGGT
GGTGGTGGAAATTCTAGACTCCATGGGATCATTAATGAAAGCTGCTGACTATGCAACGGAA
GTTTCAATATGCTAAACGGCAGATTCTGCAAGCAGCTGGTACTCTCTCTGGCGCAG
GCTAACCAACCTCCGCAAAACGGCTCTCTTACTGGTAAAG

Protein sequence (253 AA):

MRGSHHHHHGMASMTGGQQMGRDLYYDDDKDPMAQVINTNSLTLQNNLNKSQSSLSSAIERLSS
GLRINSAKDDAACQAIAANRFTSNIKGLTQASRNANDGISIAQTTEGALNEINNNLQRVRELSVQATI
PGISGGGGILDMSGTLIN
EDAAAACKSTANPLASIDSALSKVDAVRSSLGATQNRFDSAITNLGNT
VINLNSARSRIEDADYATEVSNMSKAQILQQAGTSVLAQANQVPQNVLSSLR

AB'n1-100

Nucleotide sequence (597 bp):

ATGCCGGGTTCTCATCATCATCATGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTGACGGATGACGATAAGGAATCCGATGGCACAGTCATTAAACA
AACAGCCCTCGCTGGGATCTGTGACGGATGACGATAACCTGAACAAATCTCAGTCCTACTGAGTTCC
GCTATTGAGGCTCTGCTCTGGTCTGGTCTGAGGCTGACTCTGAGGCTTCCGGTAAAC
GCTAACGACGGCATTCTTATGGCGCAGACCACTGAAGGGTGGCTGAATGAAATCAACAC
AACCTGCAACGGCTGGTGTGAGGTTGCTGAGGCTACTATCCGGGAATTCCGGTGGT
GGTGGTGGAAATTCTAGACTCCATGGGATCATTAATGAAAGACGCTGGCGCAGCCAAAG
AAAAGTACCCCTAACCAACTGGCTTCAATGATTTGCTCTGCTAAAGTGGACCGAGTT
GTTCTCTCTGGGGCAATTCAAACCAACCTTTGATTGACCCATTACCAACCTTTAG

Protein sequence (198 AA):

MRGSHHHHHGMASMTGGQQMGRDLYYDDDKDPMAQVINTNSLTLQNNLNKSQSSLSSAIERLSS
GLRINSAKDDAACQAIAANRFTSNIKGLTQASRNANDGISIAQTTEGALNEINNNLQRVRELSVQATI
PGISGGGGILDMSGTLIN
EDAAAACKSTANPLASIDSALSKVDAVRSSLGATQNRFDSAITNL

Fig. 38 - Continued

AA'n1-70

Nucleotide sequence (672 bp):

```

ATGGGGGGTTCTCATCATCATCATCATCAGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTACGACGATGACGATAAGGATCCGATGGCACAAAGTCATTAAACA
AACAGCCTGTCGTGTTGACCCAGAATAACCTGACAAATCTCAGTCCTCACTGAATTCC
GCTATTGAGCGTCCTGCTCTGGCTATCAACAGCGGAAAGACGATCGGGTAGGC
CAGGCATTCGCTAACCGCTTCACTTCTAATACTAAAGGCCCTGACTCAGGCTTCCCGTAAC
GCTAACGACATCCCCGAATTCCGGTGGTGGCTGGAAATTCTAGACTCCATGGCTACA
TTAACATGAAAGACGCTCCCGCAGCCAAGAAAAGTACCGCTAACCCACTGGCTTCATT
GATTCTGCATTGTCAAAGTGAACGCAGTTCTGCTCTCTGGGGCAATTCAAAACCGT
TTTGATTTCAGCCATTACCAACCTTGGCAATAACGTAACCAATCTGAACCTGGCGTAGC
CGTATCGAACGATGCTGACTATGCAACGGAAGTTCTAAATATGTCTAAAGCGCAGATCTG
CAGCAGGCTGGTACTTCGGTTCTGGCCAGGGTAACCAAGGTTCCGAAACGTCCTCTCT
TTACTGCGTTAG

```

Protein sequence (223 AA):

```

MRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPMAQVINTNSLSSLTONNLNKSQSSLSSAIERLSS
GLRINSAKDDAAGOQAIANRFTSNIKGLTQASRNANDIPGISSGGGILDSMGTLINEDEAAAKKSTA
NPLASIDSALSKVDAVRSILGAIQNRFDSTAINTLGNVTINLNSARSRIEDADYATEVSNMSKAQILQ
QAGTSVLAQANQVPQNVLSLLR

```

AB'n1-70

Nucleotide sequence (507 bp):

```

ATGGGGGGTTCTCATCATCATCATCAGGTATGGCTAGCATGACTGGTGGACAGCAA
ATGGGTGGGATCTGTACGACGATGACGATAAGGATCCGATGGCACAAAGTCATTAAACA
AACAGCCTGTCGTGTTGACCCAGAATAACCTGACAAATCTCAGTCCTCACTGAATTCC
GCTATTGAGCGTCCTGCTCTGGCTATCAACAGCGGAAAGACGATCGGGTAGGC
CAGGCATTCGCTAACCGCTTCACTTCTAATACTCAACGGCTGACTCAGGCTTCCCGTAAC
GCTAACGACATCCCCGAATTCCGGTGGTGGCTGGAAATTCTAGACTCCATGGCTACA
TTAACATGAAAGACGCTCCCGCAGCCAAGAAAAGTACCGCTAACCCACTGGCTTCATT
GATTCTGCATTGTCAAAGTGGACCCAGTTCTGCTCTCTGGGGCAATTCAAAACCGT
TTTGATTTCAGCCATTACCAACCTTGTAG

```

Protein sequence (168 AA):

```

MRGSHHHHHGMASMTGGQQMGRDLYDDDDKDPMAQVINTNSLSSLTONNLNKSQSSLSSAIERLSS
GLRINSAKDDAAGOQAIANRFTSNIKGLTQASRNANDIPGISSGGGILDSMGTLINEDEAAAKKSTA
NPLASIDSALSKVDAVRSSLGAIQNRFDSTAINTL

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FLAGELLIN RELATED POLYPEPTIDES AND USES THEREOF

CROSS REFERENCE TO RELATED APPLICATIONS

This application is a continuation of U.S. patent application Ser. No. 14/559,669, filed on Dec. 3, 2014, now U.S. Pat. No. 9,139,623, which is a continuation of U.S. patent application Ser. No. 13/110,704, filed on May 18, 2011, now U.S. Pat. No. 8,932,609, which is a divisional of U.S. patent application Ser. No. 11/722,682, filed on May 2, 2008, now U.S. Pat. No. 8,007,812, which is the national stage of International Application No. PCT/US2005/046485, filed on Dec. 22, 2005, which claims the benefit of U.S. Provisional Patent Application No. 60/639,826, filed Dec. 22, 2004, the contents of all of which are incorporated herein by reference.

FIELD OF THE INVENTION

This invention relates to the use of flagellin related polypeptides to protect mammals from the effects of apoptosis. More specifically, this invention relates to the use of flagellin related polypeptides to protect mammals from exposure to stress, such as radiation and cancer treatments.

REFERENCE TO THE SEQUENCE LISTING

Reference is made to the sequence listing submitted via EFS-Web, which consists of a file named, "CLE-003D5-SequenceListing.txt" (135 KB), created on Aug. 17, 2015, the contents of which are incorporated herein by reference.

BACKGROUND OF THE INVENTION

The progression from normal cells to tumor cells involves a loss of negative mechanisms of growth regulation, including resistance to growth inhibitory stimuli and a lack of dependence on growth factors and hormones. Traditional cancer treatments that are based on radiation or cytotoxic drugs rely on the differences in growth control of normal and malignant cells. Traditional cancer treatments subject cells to severe genotoxic stress. Under these conditions, the majority of normal cells become arrested and therefore saved, while tumor cells continue to divide and die.

However, the nature of conventional cancer treatment strategy is such that normal rapidly dividing or apoptosis-prone tissues are at risk. Damage to these normal rapidly dividing cells causes the well-known side effects of cancer treatment (sensitive tissues: hematopoiesis, small intestine, hair follicles). The natural sensitivity of such tissues is complicated by the fact that cancer cells frequently acquire defects in suicidal (apoptotic) machinery and those therapeutic procedures that cause death in normal sensitive tissues may not be that damaging to cancer cells. Conventional attempts to minimize the side effects of cancer therapies are based on (a) making tumor cells more susceptible to treatment, (b) making cancer therapies more specific for tumor cells, or (c) promoting regeneration of normal tissue after treatment (e.g., erythropoietin, GM-CSF, and KGF).

There continues to be a need for therapeutic agents to mitigate the side effects associated with chemotherapy and radiation therapy in the treatment of cancer. This invention fulfills these needs and provides other related advantages.

SUMMARY OF THE INVENTION

A method of protecting a mammal from one or more treatments or conditions that trigger apoptosis comprising admin-

istering to said patient a composition comprising a pharmaceutically effective amount of flagellin. The flagellin may comprise SEQ ID NO: 1 or a fragment, variant, analog, homolog, derivative of SEQ ID NO: 1, or combination thereof. The flagellin may induce TLR-5 mediated activity.

The flagellin may be at least 30% identical to amino acids 1-174 and 418-505 of SEQ ID NO: 1. The flagellin may comprise at least 10 conserved amino acids at positions selected from the group consisting of 89, 90, 91, 95, 98, 101, 115, 422, 423, 426, 431, 436 and 452. The flagellin may comprise the sequence of SEQ ID NOS: 1, 8, 10, 12, 30, 32, 34, 36, 38, 40, 43, 44, 46, 48, 50 and 52.

The flagellin may be used to treat a mammal undergoing cancer treatment, which may be chemotherapy or radiation therapy. The flagellin may be used to treat a mammal exposed to radiation. The flagellin may be administered in combination with a radioprotectant. The flagellin may be used to treat a mammal from wounding, poisoning, bacterial infection, viral infection and temperature shock. The flagellin may be used to protect from apoptosis in tissues including the GI tract, lungs, kidneys, liver, cardiovascular system, blood vessel endothelium, central and peripheral neural system, hematopoietic progenitor cells, immune system, and hair follicles. The flagellin may also be used to prevent sepsis in the mammal.

This invention also relates to a method of treating a mammal suffering from a constitutively active NF- κ B cancer comprising administering to the mammal a composition comprising a pharmaceutically acceptable amount of an agent which induces NF- κ B. The agent may be flagellin. The agent may be administered prior to, together with, or after a treatment for the cancer. The treatment may be chemotherapy or radiation therapy.

This invention also relates to a method of treating a mammal suffering from damage to normal tissue attributable to treatment of a cancer comprising administering to the mammal a composition comprising a pharmaceutically acceptable amount of an agent which induces NF- κ B. The agent may be flagellin. The agent may be administered prior to, together with, or after a treatment for the cancer. The treatment may be chemotherapy or radiation therapy.

This invention also relates to a method of treating a mammal suffering from damage to normal tissue attributable to stress, comprising administering to the mammal a composition comprising a pharmaceutically acceptable amount of an agent which induces NF- κ B. The agent may be flagellin. The agent may be administered prior to, together with, or after a treatment for a disease suffered by the mammal.

This invention also relates to a method of modulating cell aging in a mammal, comprising administering to the mammal a composition comprising a pharmaceutically acceptable amount of an agent which induces NF- κ B. The agent may be flagellin. The agent may be administered prior to, together with, or after a treatment for a disease suffered by the mammal.

This invention also relates to a pharmaceutical composition comprising an agent which induces NF- κ B activity, a chemotherapeutic drug, and optionally a pharmaceutically acceptable adjuvant, diluent, or carrier. The agent may be flagellin.

This invention also relates to a method of screening for an inducer of NF- κ B comprising adding a suspected inducer to an NF- κ B activated expression system, and separately adding a control to an NF- κ B activated expression system, whereby an inducer of NF- κ B is identified by the ability to increase the level of NF- κ B activated expression.

This invention also relates to a method of protecting a mammal from the effects of radiation comprising administering to said mammal a composition comprising a pharmaceutically effective amount of an agent which induces NF- κ B. The agent may be flagellin, which may be derived from a species of *Salmonella*. The composition may be administered in combination with a radioprotectant. The radioprotectant may be an antioxidant, which may be amifostine or vitamine E. The radioprotectant may also be a cytokine, which may be stem cell factor.

This invention relates to a method of protecting a patient from one or more treatments or conditions that trigger apoptosis comprising administering to said patient a composition comprising a pharmaceutically effective amount of an agent which induces NF- κ B. The agent may be flagellin, which may be derived from a species of *Salmonella*. The treatment may be a cancer treatment, which may be chemotherapy or radiation therapy. The condition may be a stress, which may be radiation, wounding, poisoning, infection and temperature shock.

This invention also relates to a method of screening for a modulator of apoptosis comprising adding a suspected modulator to a cell-based apoptosis system, and separately adding a control to a cell-based apoptosis system, whereby a modulator of apoptosis is identified by the ability to alter the rate of apoptosis, wherein the suspected modulator is derived from a mammalian parasite or symbiont.

This invention also relates to a method of screening for a modulator of NF- κ B comprising adding a suspected modulator to an NF- κ B activated expression system, and separately adding a control to an NF- κ B activated expression system, whereby a modulator of NF- κ B is identified by the ability to alter the rate of NF- κ B activated expression, wherein the suspected modulator is derived from a mammalian parasite. The parasite may be of a species including, but not limited to, *Salmonella*, *Mycoplasma*, and *Chlamydia*.

This invention also relates to a modulator identified by any of the screening methods described herein. This invention also relates to a composition comprising a modulator described herein. The composition may be a pharmaceutical composition comprising a pharmaceutically acceptable amount of a modulator described herein.

This invention also relates to a method of treating cancer comprising administering to a subject in need of such treatment a pharmaceutical composition comprising a modulator that enhances apoptosis.

This invention also relates to a method of protecting a patient from one or more treatments that trigger apoptosis comprising administering to said patient a pharmaceutical composition comprising a modulator that inhibits apoptosis. The one or more treatments may be a cancer treatment. The cancer treatment may be chemotherapy or radiation therapy.

BRIEF DESCRIPTION OF THE DRAWINGS

FIG. 1 demonstrates that p53 deficiency accelerated development of GI syndrome in mice. Panel A: I.P. injection of PFT α (10 mg/kg) protects C57Bl/6J mice (if not indicated otherwise, here and below 6-8 weeks old males were used) from a single 9 Gy dose of gamma radiation and a fractionated cumulative radiation dose 12.5 Gy (5 \times 2.5 Gy). PFT α has no effect on survival of mice treated with single 12.5 and 25 Gy doses of IR: (results of representative experiments are shown; Shepherd 4000 Ci Cesium 137 source at a dose rate of 4 Gy per minute was used). Panel B: Wild-type and p53-null C57Bl/6J mice differ in their relative sensitivity to low (10 Gy) and high (15 Gy) doses of gamma radiation: wild-type

mice were more sensitive to 10 Gy but more resistant to 15 Gy as compared to p53-null mice. Panel C: Mice treated with 11 Gy of total body gamma irradiation were injected 12 h later with 1.5×10^7 bone marrow cells from wild type or p53-null syngeneic C57Bl/6J mice. (This dose causes 100% lethality in nonreconstituted controls group of mice). Two months later, after complete recovery of hematopoiesis, animals were treated with 15 Gy of total body gamma radiation and showed no difference in death rates between the two groups differing 10 in the p53 status of their bone marrow. Panel D: Comparison of dynamics of injury to small intestines of wild-type and p53-null mice at the indicated time points after 15 Gy of gamma radiation indicates accelerated damage in p53-null mice (haematoxylin-eosin stained paraffin sections; magnification \times 125). 24 h panels include images of TUNEL staining if sections of crypts: massive apoptosis is evident in wild type but not in p53-deficient epithelium.

FIG. 2 demonstrates the dynamics of cell proliferation and survival in small intestine of wild type and p53-null mice. 20 Panel A: Comparison of proliferation rates in intestines of wild-type and p53 null mice after treatment with IR. (Left) Autoradiographs of whole-body sections (1.7 \times magnification) of 4-week-old wild-type and p53 null mice injected intraperitoneally with 14 C-thymidine (10 μ Ci per animal) treated or untreated with 15 Gy of gamma radiation. Arrows point at intestines. (Right) Comparison of BrdU incorporation in small intestine of wild-type and p53-null mice at different time points after 15 Gy of gamma radiation. BrdU (50 mg/kg) was injected 2 h before sacrificing mice followed 30 by immunostaining. Fragments of 96 h panels are shown at higher magnification (\times 400). Panel B: Comparison of the number of BrdU positive cells/crypt in small intestine of wild-type and p53-null mice at different time points after 15 Gy of gamma radiation. Three animals were analyzed for each time point, five ileum cross sections were prepared from each animal and analyzed microscopically to estimate the number of crypts and villi. Numbers of BrdU-positive cells in the crypts were counted in 5 random fields under 200 \times magnification (100-30 crypts) and the average number of BrdU-positive cells was plotted. Panel C: Tracing the number and position of BrdU-labeled cells in small intestine of wild type and p53-null mice during different time points after 15 Gy of gamma radiation. BrdU was injected 30 min. before irradiation and mice were sacrificed at the indicated time points. 40 Accelerated migration from crypts to villi followed by rapid elimination of labeled cells was observed in p53-null mice.

FIG. 3 demonstrates that recombinant flagellin is capable of NF- κ B activation.

FIG. 4 shows a representative experiment testing the ability 50 of flagellin to protect mice from radiation. C56BL6 mice (6 week old males, 10 animals per group) were injected i.v. with 2.014 (0.1 mg/kg) or 514 (0.25 mg/kg) of flagellin in PBS. Four hours later, mice were irradiated with 15 Gy and mouse survival was monitored daily.

FIG. 5 shows histological sections (HE stained) of small 55 intestinal epithelium of mice that were treated with 15 Gy of gamma radiation with or without i.v. injection of 0.25 mg/kg of flagellin. Complete destruction of crypts and villi in control mouse contrasts with close to normal morphology of tissue from flagellin-treated animal.

FIG. 6 shows the effect of flagellin on mouse sensitivity to 10 Gy of total body gamma radiation.

FIG. 7 shows the effect of flagellin injected i.v. at indicated 60 times before irradiation on mouse sensitivity to 13 Gy (left) and 10 Gy (right) of total body gamma radiation.

FIG. 8 shows the effect of flagellin on mouse sensitivity to 10, 13 and 15 Gy of total body gamma radiation.

FIG. 9 shows the domain structure of bacterial flagellin. The Ca backbone trace, hydrophobic core distribution and structural information of F41. Four distinct hydrophobic cores that define domains D1, D2a, D2b and D3. All the hydrophobic side-chain atoms are displayed with the Ca backbone. Side-chain atoms are color coded: Ala, yellow; Leu, Ile or Val, orange; Phe and Tyr, purple (carbon atoms) and red (oxygen atoms). c, Position and region of various structural features in the amino-acid sequence of flagellin. Shown are, from top to bottom: the F41 fragment in blue; three b-folium folds in brown; the secondary structure distribution with a-helix in yellow, b-structure in green, and b-turn in purple; tic mark at every 50th residue in blue; domains D0, D1, D2 and D3; the axial subunit contact region within the proto-element in cyan; the well-conserved amino-acid sequence in red and variable region in violet; point mutations in F41 that produce the elements of different supercoils. Letters at the bottom indicate the morphology of mutant elements: L (D107E, R124A, R124S, G426A), L-type straight; R (A449V), R-type straight; C (D313Y, A414V, A427V, N433D), curly33.

FIG. 10 shows a schematic of *Salmonella* flagellin domains, its fragments, and its interaction with TLR5. Dark bars denote regions of the flagellin gene used to construct fragments comprising A, B, C, A' and B'.

FIG. 11 shows soluble flagellin (FliC), and two fragments (AA' and BB') after fractionation by SDS-PAGE, with molecular weight markers listed to the left.

FIG. 12 shows induction of NF- κ B nuclear translocation by *Salmonella* flagellin (FliC) and flagellin fragments.

FIG. 13 shows activation of NF- κ B-regulated luciferase reporter construct by flagellin and flagellin fragments in H116 cells. Concentrations of proteins are given in μ g/ml.

FIG. 14 shows NF- κ B DNA binding in HT29 human colon cancer cells induced by flagellin and flagellin fragments.

FIG. 15 shows the activation of a NF- κ B reporter in HCT116 reporter cells by full-length flagellin and flagellin fragments.

FIG. 16 shows a comparison of the radioprotective properties of flagellin (FliC) and fragments AA' and BB'.

FIG. 17 shows that the AA' fragment protects intestinal epithelium from degeneration caused by radiation. A: Histological sections (hematoxylin and eosin-stained) of small intestinal epithelium of mice 5 days after 14 Gy irradiation are shown. B: Treatment with the AA' fragment prevents apoptosis ongoing 5 hours after irradiation in endothelial cells of villi (detected by immunostaining for endothelial marker CD31 and marked by arrows), as determined by TUNEL assay. C: Histological sections of skin of mice 5 days after 14 Gy of gamma irradiation demonstrate the protective effect of the AA' fragment for sebaceous glands (red arrows).

FIG. 18 shows that the AA' fragment provides partial protection and delays death of mice after supralethal irradiation with 17 and 20 Gy total-body gamma radiation.

FIG. 19 shows anti-flagellin antibody titers induced in mice after 21 and 28 days by flagellin and AA'. For individual mice, the averages of two measurements are shown. Mice were injected with: Fl: flagellin; or AA'. 21d and 28 ds—mice injected with first dose 21 and 28 days before second, respectively. PBS: saline buffer (no serum) control; blank: empty well reading control.

FIG. 20 shows anti-flagellin antibody titers induced in mice after 21 and 28 days by flagellin and AA'. For individual mice, the averages of two measurements are shown. Mice were injected with: Fl: flagellin; or AA'. 21 d and 28 ds—mice

injected with first dose 21 and 28 days before second, respectively. PBS: saline buffer (no serum) control; blank: empty well reading control.

FIG. 21 shows that flagellin fragment AA' protects mice from multiple successive doses of gamma-irradiation. Arrows denote radiation treatments (days 1-4).

FIG. 22 shows the effect of AA' on tumor sensitivity to radiation treatment. Left Panel: NIH3T3-derived sarcoma cells were injected s.c. in NIH-Swiss mice. When tumors reached 7-10 mm in diameter, mice received three 4.3 Gy doses of total body irradiation, with or without pretreatment with AA'. The dynamics of tumor growth after radiation treatment is displayed. U/t: untreated; AA': AA' with no irradiation; 3x4 Gy: irradiation only; 3x4 Gy+AA':AA' and irradiation. (The shape of curves reflects slow growth of tumors that is a characteristic of this model). Results are displayed as relative tumor volumes normalized to tumor volume measured at day 7 after last irradiation. Right Panel: The experiment was done in the same way with another syngeneic mouse tumor model: B16 melanoma (C57BL6 background). Treatment was applied when tumors reached 4-5 mm in diameter and involved three subsequent 4 Gy doses of total body gamma radiation applied with or without pretreatment with AA' (30 min. before irradiation, 5 μ g/mouse).

FIG. 23 shows the influence of NS398 on the radioprotection of LPS and AA' in mice after 13 Gy of total-body gamma irradiation.

FIGS. 24A and 24B show a comparison of amino acid sequences of the conserved amino (FIG. 24A) and carboxy (FIG. 24B) terminus from 21 species of bacteria. The 13 conserved amino acids important for TLR5 activity are shown with shading. The amino acid sequences are identified by their accession numbers from TrEMBL (first letter=Q) or Swiss-Prot (first letter=P). The amino terminus sequences have SEQ ID NOs: 1-21, respectively, for each of the 21 bacterial species, and the carboxy terminus sequences have SEQ ID NOs: 22-42, respectively.

FIGS. 25A-D show results of a BLAST search using SEQ ID NO: 1 as the query sequence. The parameters used in all searches was as follows: expected value cutoff=10, matrix=BLOSUM62, gap penalties of existence=11 and extension=1, filtering=none. FIG. 25A: NR_Bacteria (Protein-Protein); FIG. 25B: NR_Bacteria (Protein-DNA); FIG. 25C: Bacterial Genomes (Protein-Protein); FIG. 25D: Bacterial Genomes (Protein-DNA).

FIG. 26 shows the percentage identities of the amino- and carboxy-terminus of the homologs shown in FIG. 24 compared to SEQ ID NO: 1, as shown in BLAST results using the same search parameters as listed for FIGS. 25A-D.

FIG. 27 demonstrates that AA' mediates rescue of multiple mouse strains after 10 Gy total-body γ -IR. Cone heights represent fractions of survivors.

FIG. 28 demonstrates the pharmacokinetics of AA' after intravenous (i.v.), subcutaneous (s.c.), intraperitoneal (i.p.) or intramuscular (i.m.) injection.

FIG. 29 demonstrates the extended pharmacokinetics of AA' after intramuscular (i.m.) injection.

FIG. 30 demonstrates the influence of AA' on gamma-irradiation induced cell death and growth inhibition in A549 cells.

FIG. 31 demonstrates the influence of AA' on gamma-irradiation induced cell death and growth inhibition in multiple cell lines.

FIG. 32 demonstrates the influence of irradiation and AA' on BrdU incorporation in small intestinal crypts of NIH-Swiss mice. A comparison of BrdU incorporation in small intestine of control and AA'-treated NIH-Swiss mice, with and

without 15 Gy of gamma radiation is shown. BrdU (50 mg/kg) was injected 1.5 h before sacrificing mice and immunostaining was done as previously described (Watson A J & Pritchard D M., *Am J Physiol Gastrointest Liver Physiol.* 2000 January; 278(1):G 1-5). Red channel of the image is shown (positive signal is bright white on the dark background).

FIG. 33 demonstrates the duration of AA'-mediated growth arrest and reduced BrdU incorporation in small intestine of mice. BrdU (50 mg/kg) was injected in Balb/c mice i.p., 1 or 4 hrs after CBLB502 (AA') injection. Samples of small intestine were obtained 1.5 hrs after BrdU injection. Immunostaining was done as previously described (Watson A J & Pritchard D M., *Am J Physiol Gastrointest Liver Physiol.* 2000 January; 278(1):G 1-5). Inverted image is shown (positive signal is dark on the light background).

FIG. 34 demonstrates the influence of AA' on BrdU incorporation in colonic crypts of NIH-Swiss mice. BrdU (50 mg/kg) was injected in NIH-Swiss mice i.p., 1 hr after CBLB502 (AA') injection. Samples of colon were obtained 1.5 hrs after BrdU injection. Immunostaining was done as previously described (Watson A J & Pritchard D M., *Am J Physiol Gastrointest Liver Physiol.* 2000 January; 278(1):G 1-5). Inverted image is shown (positive signal is dark on the light background). Bottom panel shows smaller magnification/larger area of the sample.

FIG. 35 demonstrates the morphology of small intestine in TLR5 deficient MOLF/Ei and TLR5 wt NIH-Swiss mice after treatment with AA'.

FIG. 36 depicts flagellin derivatives. The domain structure and approximate boundaries (amino acid coordinates) of selected flagellin derivatives (listed on the right). FliC flagellin of *Salmonella dublin* is encoded within 505 amino acids (aa).

FIG. 37 shows the testing of additional flagellin derivatives tested for NF-κB stimulating activity.

FIG. 38 shows the nucleotide and amino acid sequence for the following flagellin variants: AA' (SEQ ID NO: 7-8), AB' (SEQ ID NO: 9-10), BA' (SEQ ID NO: 11-12), BB' (SEQ ID NO: 13-14), CA' (SEQ ID NO: 15-16), CB' (SEQ ID NO: 17-18), A (SEQ ID NO: 19-20), B (SEQ ID NO: 21-22), C (SEQ ID NO: 23-24), GST-A' (SEQ ID NO: 25-26), GST-B' (SEQ ID NO: 27-28), AA'n1-170 (SEQ ID NO: 29-30), AA'n1-163 (SEQ ID NO: 33-34), AA'n54-170 (SEQ ID NO: 31-32), AA'n54-163 (SEQ ID NO: 335-36), AB'n1-170 (SEQ ID NO: 37-38), AB'n1-163 (SEQ ID NO: 39-40), AA'n1-129 (SEQ ID NO: 41-42), AA'n54-129 (SEQ ID NO: 43-44), AB'n1-129 (SEQ ID NO: 45-46), AB'n54-129 (SEQ ID NO: 47-48), AA'n1-100 (SEQ ID NO: 49-50), AB'n1-100 (SEQ ID NO: 51-52), AA'n1-70 (SEQ ID NO: 53-54) and AB'n1-70 (SEQ ID NO: 55-56). The pRSETb leader sequence is shown in Italic (leader includes Met, which is also amino acid 1 of FliC). The N terminal constant domain is underlined. The amino acid linker sequence is in Bold. The C terminal constant domain is underlined. GST, if present, is highlighted.

DETAILED DESCRIPTION

This invention is related to protecting normal cells and tissues from apoptosis caused by stresses including, but not limited to, chemotherapy, radiation therapy and radiation. There are two major mechanisms controlling apoptosis in the cell: the p53 pathway (pro-apoptotic) and the NF-κB pathway (anti-apoptotic). Both pathways are frequently deregulated in tumors: p53 is usually lost, while NF-κB becomes constitutively active. Hence, inhibition of p53 and activation of NF-κB in normal cells may protect them from death caused

by stresses, such as cancer treatment, but would not make tumor cells more resistant to treatment because they have these control mechanisms deregulated. This contradicts the conventional view on p53 and NF-κB, which are considered as targets for activation and repression, respectively.

This invention relates to inducing NF-κB activity to protect normal cells from apoptosis. By inducing NF-κB activity in a mammal, normal cells may be protected from apoptosis attributable to cellular stress, which occurs in cancer treatments and hyperthermia; exposure to harmful doses of radiation, for example, workers in nuclear power plants, the defense industry or radiopharmaceutical production, and soldiers; and cell aging. Since NF-κB is constitutively active in many tumor cells, the induction of NF-κB activity may protect normal cells from apoptosis without providing a beneficial effect to tumor cells. Once the normal cells are repaired, NF-κB activity may be restored to normal levels. NF-κB activity may be induced to protect such radiation- and chemotherapy-sensitive tissues as the hematopoietic system (including immune system), the epithelium of the gut, and hair follicles.

Inducers of NF-κB activity may also be used for several other applications. Pathological consequences and death caused by exposure of mammals to a variety of severe conditions including, but not limited to, radiation, wounding, poisoning, infection, aging, and temperature shock, may result from the activity of normal physiological mechanisms of stress response, such as induction of programmed cell death (apoptosis) or release of bioactive proteins, cytokines.

Apoptosis normally functions to "clean" tissues from wounded and genetically damaged cells, while cytokines serve to mobilize the defense system of the organism against the pathogen. However, under conditions of severe injury both stress response mechanisms can by themselves act as causes of death. For example, lethality from radiation may result from massive p53-mediated apoptosis occurring in hematopoietic, immune and digestive systems. Rational pharmacological regulation of NF-κB may increase survival under conditions of severe stress. Control over these factors may allow control of both inflammatory response and the life-death decision of cells from the injured organs. Tissues that may be protected from apoptosis by administering NF-κB inducers include, but are not limited to, the GI tract, lungs, kidneys, liver, 130504.1 cardiovascular system, blood vessel endothelium, central and peripheral neural system, hematopoietic progenitor cells, immune system, and hair follicles.

The protective role of NF-κB is mediated by transcriptional activation of multiple genes coding for: a) anti-apoptotic proteins that block both major apoptotic pathways, b) cytokines and growth factors that induce proliferation and survival of HP and other stem cells, and c) potent ROS-scavenging antioxidant proteins, such as MnSOD (SOD-2). Thus, by temporal activation of NF-κB for radioprotection, it may be possible to achieve not only suppression of apoptosis in cancer patients, but also the ability to reduce the rate of secondary cancer incidence because of simultaneous immunostimulatory effect, which, may be achieved if activation of NF-κB is reached via activation of Toll-like receptors.

Another attractive property of the NF-κB pathway as a target is its activation by numerous natural factors that can be considered as candidate radioprotectants. Among these, are multiple pathogen-associated molecular patterns (PAMPs). PAMPs are molecules that are not found in the host organism, are characteristic for large groups of pathogens, and cannot be easily mutated. They are recognized by Toll-like receptors (TLRs), the key sensor elements of innate immunity. TLRs

act as a first warning mechanism of immune system by inducing migration and activation of immune cells directly or through cytokine release. TLRs are type I membrane proteins, known to work as homo- and heterodimers. Upon ligand binding, TLRs recruit MyD88 protein, an indispensable signaling adaptor for most TLRs. The signaling cascade that follows leads to effects including (i) activation of NF- κ B pathway, and (ii) activation of MAPKs, including Jun N-terminal kinase (JNK). The activation of the NF- κ B pathway by Toll-like receptor ligands makes the ligands attractive as potential radioprotectors. Unlike cytokines, many PAMPs have little effect besides activating TLRs and thus are unlikely to produce side effects. Moreover, many PAMPs are present in humans.

Consistently with their function of immunocyte activation, all TLRs are expressed in spleen and peripheral blood leukocytes, with more TLR-specific patterns of expression in other lymphoid organs and subsets of leukocytes. However, TLRs are also expressed in other tissues and organs of the body, e.g., TLR1 is expressed ubiquitously, TLR5 is also found in GI epithelium and endothelium, while TLRs 2, 6, 7 and 8 are known to be expressed in lung.

1. Definitions

It is to be understood that the terminology used herein is for the purpose of describing particular embodiments only and is not intended to be limiting. It must be noted that, as used in the specification and the appended claims, the singular forms "a," "an" and "the" include plural referents unless the context clearly dictates otherwise.

As used herein, the terms "administer" when used to describe the dosage of an agent that induces NF- κ B activity, means a single dose or multiple doses of the agent.

As used herein, the term "analog", when used in the context of a peptide or polypeptide, means a peptide or polypeptide comprising one or more non-standard amino acids or other structural variations from the conventional set of amino acids.

As used herein, the term "antibody" means an antibody of classes IgG, IgM, IgA, IgD or IgE, or fragments, fragments or derivatives thereof, including Fab, F(ab')₂, Fd, and single chain antibodies, diabodies, bispecific antibodies, bifunctional antibodies and derivatives thereof. The antibody may be a monoclonal antibody, polyclonal antibody, affinity purified antibody, or mixtures thereof which exhibits sufficient binding specificity to a desired epitope or a sequence derived therefrom. The antibody may also be a chimeric antibody. The antibody may be derivatized by the attachment of one or more chemical, peptide, or polypeptide moieties known in the art. The antibody may be conjugated with a chemical moiety.

As used herein, "apoptosis" refers to a form of cell death that includes progressive contraction of cell volume with the preservation of the integrity of cytoplasmic organelles; condensation of chromatin (i.e., nuclear condensation), as viewed by light or electron microscopy; and/or DNA cleavage into nucleosome-sized fragments, as determined by centrifuged sedimentation assays. Cell death occurs when the membrane integrity of the cell is lost (e.g., membrane blebbing) with engulfment of intact cell fragments ("apoptotic bodies") by phagocytic cells.

As used herein, the term "cancer" means any malignant growth or tumor caused by abnormal and uncontrolled cell division; it may spread to other parts of the body through the lymphatic system or the blood stream.

As used herein, the term "cancer treatment" means any treatment for cancer known in the art including, but not limited to, chemotherapy and radiation therapy.

As used herein, the term "combination with" when used to describe administration of an agent that induces NF- κ B activity and an additional treatment means that the agent may be administered prior to, together with, after, or metronomically with the additional treatment. The term "together with," "simultaneous" or "simultaneously" as used herein, means that the additional treatment and the agent of this invention are administered within 48 hours, preferably 24 hours, more preferably 12 hours, yet more preferably 6 hours, and most preferably 3 hours or less, of each other. The term "metronomically" as used herein means the administration of the agent at times different from the additional treatment and at certain frequency relative to repeat administration and/or the additional treatment.

The agent may be administered at any point prior to the additional treatment including, but not limited to, about 48 hr, 46 hr, 44 hr, 42 hr, 40 hr, 38 hr, 36 hr, 34 hr, 32 hr, 30 hr, 28 hr, 26 hr, 24 hr, 22 hr, 20 hr, 18 hr, 16 hr, 14 hr, 12 hr, 10 hr, 8 hr, 6 hr, 4 hr, 3 hr, 2 hr, or 1 hr prior to the additional treatment.

The agent may be administered at any point after the additional treatment including, but not limited to, about 1 hr, 2 hr, 3 hr, 4 hr, 6 hr, 8 hr, 10 hr, 12 hr, 14 hr, 16 hr, 18 hr, 20 hr, 22 hr, 24 hr, 26 hr, 28 hr, 30 hr, 32 hr, 34 hr, 36 hr, 38 hr, 40 hr, 42 hr, 44 hr, 46 hr, or 48 hr after exposure.

As used herein, the term "derivative", when used in the context of a peptide or polypeptide, means a peptide or polypeptide different other than in primary structure (amino acids and amino acid analogs). By way of illustration, derivatives may differ by being glycosylated, one form of post-translational modification. For example, peptides or polypeptides may exhibit glycosylation patterns due to expression in heterologous systems. If at least one biological activity is retained, then these peptides or polypeptides are derivatives according to the invention. Other derivatives include, but are not limited to, fusion peptides or fusion polypeptides having a covalently modified N- or C-terminus, PEGylated peptides or polypeptides, peptides or polypeptides associated with lipid moieties, alkylated peptides or polypeptides, peptides or polypeptides linked via an amino acid side-chain functional group to other peptides, polypeptides or chemicals, and additional modifications as would be understood in the art.

As used herein, the term "fragment", when used in the context of a peptide or polypeptide, means a portion of a reference peptide or polypeptide.

As used herein, the term "homolog", when used in the context of a peptide or polypeptide, means a peptide or polypeptide sharing a common evolutionary ancestor.

As used herein, the term "treat" or "treating" when referring to protection of a mammal from a condition, means preventing, suppressing, repressing, or eliminating the condition. Preventing the condition involves administering a composition of this invention to a mammal prior to onset of the condition. Suppressing the condition involves administering a composition of this invention to a mammal after induction of the condition but before its clinical appearance. Repressing the condition involves administering a composition of this invention to a mammal after clinical appearance of the condition such that the condition is reduced or maintained. Eliminating the condition involves administering a composition of this invention to a mammal after clinical appearance of the condition such that the mammal no longer suffers the condition.

As used herein, the term "tumor cell" means any cell associated with a cancer.

As used herein, the term "variant", when used in the context of a peptide or polypeptide, means a peptide or polypeptide that differs in amino acid sequence by the insertion,

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deletion, or conservative substitution of amino acids, but retain at least one biological activity. Representative examples of “biological activity” include, but are not limited to, the ability to bind to TLR5 and to be bound by a specific antibody. A conservative substitution of an amino acid, i.e., replacing an amino acid with a different amino acid of similar properties (e.g., hydrophilicity, degree and distribution of charged regions) is recognized in the art as typically involving a minor change. These minor changes can be identified, in part, by considering the hydropathic index of amino acids, as understood in the art. Kyte et al., *J. Mol. Biol.* 157:105-132 (1982). The hydropathic index of an amino acid is based on a consideration of its hydrophobicity and charge. It is known in the art that amino acids of similar hydropathic indexes can be substituted and still retain protein function. In one aspect, amino acids having hydropathic indexes of ± 2 are substituted. The hydrophilicity of amino acids can also be used to reveal substitutions that would result in proteins retaining biological function. A consideration of the hydrophilicity of amino acids in the context of a peptide permits calculation of the greatest local average hydrophilicity of that peptide, a useful measure that has been reported to correlate well with antigenicity and immunogenicity. U.S. Pat. No. 4,554,101, incorporated herein by reference. Substitution of amino acids having similar hydrophilicity values can result in peptides retaining biological activity, for example immunogenicity, as is understood in the art. In one aspect, substitutions are performed with amino acids having hydrophilicity values within ± 2 of each other. Both the hydrophobicity index and the hydrophilicity value of amino acids are influenced by the particular side chain of that amino acid. Consistent with that observation, amino acid substitutions that are compatible with biological function are understood to depend on the relative similarity of the amino acids, and particularly the side chains of those amino acids, as revealed by the hydrophobicity, hydrophilicity, charge, size, and other properties.

2. Methods of Treatment

a. Constitutively Active NF- κ B Tumor

This invention relates to a method of treating a mammal suffering from a constitutively active NF- κ B cancer comprising administering to the mammal a composition comprising a therapeutically effective amount of an agent that induces NF- κ B activity. The agent that induces NF- κ B activity may be administered in combination with a cancer treatment, such as chemotherapy and radiation therapy.

The cancer treatment may comprise administration of a cytotoxic agent or cytostatic agent, or combination thereof. Cytotoxic agents prevent cancer cells from multiplying by: (1) interfering with the cell’s ability to replicate DNA and (2) inducing cell death and/or apoptosis in the cancer cells. Cytostatic agents act via modulating, interfering or inhibiting the processes of cellular signal transduction which regulate cell proliferation and sometimes at low continuous levels.

Classes of compounds that may be used as cytotoxic agents include, but are not limited to, the following: alkylating agents (including, without limitation, nitrogen mustards, ethylenimine derivatives, alkyl sulfonates, nitrosoureas and triazenes): uracil mustard, chloromethine, cyclophosphamide (Cytoxan®), ifosfamide, melphalan, chlorambucil, pipobroman, triethylenemelamine, triethylenethiophosphoramide, busulfan, carmustine, lomustine, streptozocin, dacarbazine, and temozolomide; antimetabolites (including, without limitation, folic acid antagonists, pyrimidine analogs, purine analogs and adenosine deaminase inhibitors): methotrexate, 5-fluorouracil, flouxuridine, cytarabine, 6-mercaptopurine,

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6-thioguanine, fludarabine phosphate, pentostatine, and gemcitabine; natural products and their derivatives (for example, vinca alkaloids, antitumor antibiotics, enzymes, lymphokines and epipodophyllotoxins): vinblastine, vincristine, vindesine, bleomycin, dactinomycin, daunorubicin, doxorubicin, epirubicin, idarubicin, ara-c, paclitaxel (paclitaxel is commercially available as Taxol®), mithramycin, deoxycoformycin, mitomycin-c, 1-asparaginase, interferons (preferably IFN- α), etoposide, and teniposide. Other proliferative cytotoxic agents are navelbine, CPT-11, anastrazole, letrozole, capecitabine, relaxafine, cyclophosphamide, ifosamide, and droloxafine.

Microtubule affecting agents interfere with cellular mitosis and are well known in the art for their cytotoxic activity. Microtubule affecting agents useful in the invention include, but are not limited to, allocolchicine (NSC 406042), halichondrin B (NSC 609395), colchicine (NSC 757), colchicine derivatives (e.g., NSC 33410), dolastatin 10 (NSC 376128), maytansine (NSC 153858), rhizoxin (NSC 332598), paclitaxel (Taxol®, NSC 125973), Taxol® derivatives (e.g., derivatives (e.g., NSC 608832), thiocolchicine (NSC 361792), trityl cysteine (NSC 83265), vinblastine sulfate (NSC 49842), vincristine sulfate (NSC 67574), natural and synthetic epithilones including but not limited to epithilone A, epithilone B, and discodermolide (see Service, (1996) *Science*, 274:2009) estramustine, nocodazole, MAP4, and the like. Examples of such agents are also described in Bulinski (1997) *J. Cell Sci.* 110:3055-3064; Panda (1997) *Proc. Natl. Acad. Sci. USA* 94:10560-10564; Muhlradi (1997) *Cancer Res.* 57:3344-3346; Nicolaou (1997) *Nature* 387:268-272; Vasquez (1997) *Mol. Biol. Cell.* 8:973-985; and Panda (1996) *J. Biol. Chem.* 271:29807-29812.

Also suitable are cytotoxic agents such as epidophyllotoxin; an antineoplastic enzyme; a topoisomerase inhibitor; procarbazine; mitoxantrone; platinum coordination complexes such as cis-platin and carboplatin; biological response modifiers; growth inhibitors; antihormonal therapeutic agents; leucovorin; tegafur; and haematopoietic growth factors.

Cytostatic agents that may be used include, but are not limited to, hormones and steroids (including synthetic analogs): 17 α -ethynodiol, diethylstilbestrol, testosterone, prednisone, fluoxymesterone, dromostanolone propionate, testolactone, megestrolacetate, methylprednisolone, methyltestosterone, prednisolone, triamcinolone, hlorotriานisene, hydroxyprogesterone, aminoglutethimide, estramustine, medroxyprogesteroneacetate, leuprolide, flutamide, toremifene, zoladex.

Other cytostatic agents are antiangiogenics such as matrix metalloproteinase inhibitors, and other VEGF inhibitors, such as anti-VEGF antibodies and small molecules such as ZD6474 and SU6668 are also included. Anti-Her2 antibodies from Genetech may also be utilized. A suitable EGFR inhibitor is EKB-569 (an irreversible inhibitor). Also included are Imclone antibody C225 immunospecific for the EGFR, and src inhibitors.

Also suitable for use as a cytostatic agent is Casodex® (bicalutamide, Astra Zeneca) which renders androgen-dependent carcinomas non-proliferative. Yet another example of a cytostatic agent is the antiestrogen Tamoxifen® which inhibits the proliferation or growth of estrogen dependent breast cancer. Inhibitors of the transduction of cellular proliferative signals are cytostatic agents. Representative examples include epidermal growth factor inhibitors, Her-2 inhibitors, MEK-1 kinase inhibitors, MAPK kinase inhibitors, PI3 inhibitors, Src kinase inhibitors, and PDGF inhibitors.

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A variety of cancers may be treated according to this invention including, but not limited to, the following: carcinoma including that of the bladder (including accelerated and metastatic bladder cancer), breast, colon (including colorectal cancer), kidney, liver, lung (including small and non-small cell lung cancer and lung adenocarcinoma), ovary, prostate, testes, genitourinary tract, lymphatic system, rectum, larynx, pancreas (including exocrine pancreatic carcinoma), esophagus, stomach, gall bladder, cervix, thyroid, and skin (including squamous cell carcinoma); hematopoietic tumors of lymphoid lineage including leukemia, acute lymphocytic leukemia, acute lymphoblastic leukemia, B-cell lymphoma, T-cell lymphoma, Hodgkins lymphoma, non-Hodgkins lymphoma, hairy cell lymphoma, histiocytic lymphoma, and Burkitts lymphoma; hematopoietic tumors of myeloid lineage including acute and chronic myelogenous leukemias, myelodysplastic syndrome, myeloid leukemia, and promyelocytic leukemia; tumors of the central and peripheral nervous system including astrocytoma, neuroblastoma, glioma, and schwannomas; tumors of mesenchymal origin including fibrosarcoma, rhabdomyosarcoma, and osteosarcoma; and other tumors including melanoma, xeroderma pigmentosum, keratoactanthoma, seminoma, thyroid follicular cancer, teratocarcinoma, and cancers of the gastrointestinal tract or the abdominopelvic cavity.

b. Treatment of Side Effects from Cancer Treatment

This invention also relates to a method of treating a mammal suffering from damage to normal tissue attributable to treatment of a constitutively active NF- κ B cancer, comprising administering to the mammal a composition comprising a therapeutically effective amount of an agent that induces NF- κ B activity. The agent that induces NF- κ B activity may be administered in combination with a cancer treatment described above.

c. Modulation of Cell Aging

This invention also relates to a method of modulating cell aging in a mammal, comprising administering to the mammal a therapeutically effective amount of an agent that induces NF- κ B activity. The agent that induces NF- κ B activity may be administered in combination with other treatments.

d. Treatment of Stress

This invention also relates to a method of treating a mammal suffering from damage to normal tissue attributable to stress, comprising administering to the mammal a composition comprising a therapeutically effective amount of an agent that induces NF- κ B activity. The agent that induces NF- κ B activity may be administered in combination with other treatments. The stress may be attributable to any source including, but not limited to, radiation, wounding, poisoning, infection, and temperature shock.

e. Radiation

This invention is also related to the protection of cells from the effects of exposure to radiation. Injury and death of normal cells from ionizing radiation is a combination of direct radiation-induced damage to the exposed cells and an active genetically programmed cell reaction to radiation-induced stress resulting in suicidal death or apoptosis. Apoptosis plays a key role in massive cell loss occurring in several radiosensitive organs (i.e., hematopoietic and immune systems, epithelium of digestive tract, etc.), the failure of which determines general radiosensitivity of the organism.

Exposure to ionizing radiation (IR) may be short- or long-term, it may be applied as a single or multiple doses, to the whole body or locally. Thus, nuclear accidents or military attacks may involve exposure to a single high dose of whole body irradiation (sometimes followed by a long-term poisoning with radioactive isotopes). The same is true (with strict

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control of the applied dose) for pretreatment of patients for bone marrow transplantation when it is necessary to prepare hematopoietic organs for donor's bone marrow by "cleaning" them from the host blood precursors. Cancer treatment may involve multiple doses of local irradiation that greatly exceeds lethal dose if it were applied as a total body irradiation. Poisoning or treatment with radioactive isotopes results in a long-term local exposure to radiation of targeted organs (e.g., thyroid gland in the case of inhalation of ^{125}I). Finally, there are many physical forms of ionizing radiation differing significantly in the severity of biological effects.

At the molecular and cellular level, radiation particles are able to produce breakage and cross-linking in the DNA, proteins, cell membranes and other macromolecular structures. Ionizing radiation also induces the secondary damage to the cellular components by giving rise to the free radicals and reactive oxygen species (ROS). Multiple repair systems counteract this damage, such as several DNA repair pathways that restore the integrity and fidelity of the DNA, and antioxidant chemicals and enzymes that scavenge the free radicals and ROS and reduce the oxidized proteins and lipids. Cellular checkpoint systems detect the DNA defects and delay cell cycle progression until damage is repaired or decision to commit cell to growth arrest or programmed cell death (apoptosis) is reached

Radiation can cause damage to mammalian organism ranging from mild mutagenic and carcinogenic effects of low doses to almost instant killing by high doses. Overall radiosensitivity of the organism is determined by pathological alterations developed in several sensitive tissues that include hematopoietic system, reproductive system and different epithelia with high rate of cell turnover.

The acute pathological outcome of gamma irradiation leading to death is different for different doses and is determined by the failure of certain organs that define the threshold of the organism's sensitivity to each particular dose. Thus, lethality at lower doses occurs from bone marrow aplasia, while moderate doses kill faster by inducing gastrointestinal (GI) syndrome. Very high doses of radiation can cause almost instant death eliciting neuronal degeneration.

Organisms that survive a period of acute toxicity of radiation can suffer from long-term remote consequences that include radiation-induced carcinogenesis and fibrosis developing in exposed organs (e.g., kidney, liver or lungs) months and years after irradiation.

Cellular DNA is the major target of IR causing a variety of types of DNA damage (genotoxic stress) by direct and indirect (free radical-based) mechanisms. All organisms maintain DNA repair system capable of effective recovery of radiation-damaged DNA; however, errors in the DNA repair process may lead to mutations.

Tumors are generally more sensitive to gamma radiation and can be treated with multiple local doses that cause relatively low damage to normal tissue. Nevertheless, in some instances, damage of normal tissues is a limiting factor in application of gamma radiation for cancer treatment. The use of gamma-irradiation during cancer therapy by conventional, three-dimensional conformal or even more focused Beam-Cath delivery has also dose-limiting toxicities caused by cumulative effect of irradiation and inducing the damage of the stem cells of rapidly renewing normal tissues, such as bone marrow and gastrointestinal (GI) tract.

At high doses, radiation-induced lethality is associated with so-called hematopoietic and gastrointestinal radiation syndromes. Hematopoietic syndrome is characterized by loss of hematopoietic cells and their progenitors making it impossible to regenerate blood and lymphoid system. The death

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usually occurs as a consequence of infection (result of immunosuppression), hemorrhage and/or anemia. GI syndrome is caused by massive cell death in the intestinal epithelium, predominantly in the small intestine, followed by disintegration of intestinal wall and death from bacteraemia and sepsis. Hematopoietic syndrome usually prevails at the lower doses of radiation and leads to a more delayed death than GI syndrome.

In the past, radioprotectants were typically antioxidants—both synthetic and natural. More recently, cytokines and growth factors have been added to the list of radioprotectants. The mechanism of their radioprotection is considered to be a result of a facilitating effect on regeneration of sensitive tissues. There is no clear functional distinction between both groups of radioprotectants, however, since some cytokines induce the expression of cellular antioxidant proteins, such as manganese superoxide dismutase (MnSOD) and metallothionein.

The measure of protection for a particular agent is expressed by dose modification factor (DMF or DRF). DMF is determined by irradiating the radioprotector treated subject and untreated control subjects with a range of radiation doses and then comparing the survival or some other endpoints. DMF is commonly calculated for 30-day survival (LD50/30 drug-treated divided by LD50/30 vehicle-treated) and quantifies the protection of the hematopoietic system. In order to estimate gastrointestinal system protection, LD50 and DMF are calculated for 6- or 7-day survival. DMF values provided herein are 30-day unless indicated otherwise.

As shown below, inducers of NF- κ B possess strong pro-survival activity at the cellular level and on the organism as a whole. In response to super-lethal doses of radiation, inducers of NF- κ B inhibit both gastrointestinal and hematopoietic syndromes, which are the major causes of death from acute radiation exposure. As a result of these properties, inducers of NF- κ B may be used to treat the effects of natural radiation events and nuclear accidents. Moreover, since inducers of NF- κ B acts through mechanisms different from all presently known radioprotectants, they can be used in combination with other radioprotectants, thereby, dramatically increasing the scale of protection from ionizing radiation.

As opposed to conventional radioprotective agents (e.g., scavengers of free radicals), inducers of NF- κ B activity may not reduce primary radiation-mediated damage but may act against secondary events involving active cell reaction to primary damage, therefore complementing the existing lines of defense. Pifithrin-alpha, a pharmacological inhibitor of p53 (a key mediator of radiation response in mammalian cells), is an example of this new class of radioprotectants. However, the activity of p53 inhibitors is limited to protection of the hematopoietic system and has no protective effect in digestive tract (gastrointestinal syndrome), therefore, reducing therapeutic value of these compounds. Anti-apoptotic pharmaceuticals with broader range of activity are desperately needed.

Inducers of NF- κ B may be used as a radioprotective agent to extend the range of tolerable radiation doses by increasing radiosensitivity beyond the levels achievable by currently available measures (shielding and application of existing bioprotective agents) and drastically increase the chances of survival, for example, in case of onboard nuclear accidents or large-scale solar particle events. With an approximate DMF (30-day survival) greater than 1.5, the NF- κ B inducer flagellin is more effective than any currently reported natural compound.

Inducers of NF- κ B may be also useful for treating irreparable cell loss caused by low-dose irradiation, for

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example, in the central nervous system and reproductive organs. Inducers of NF- κ B may also be used during cancer chemotherapy to treat the side effects associated with chemotherapy, including alopecia.

5 In one embodiment, a mammal is treated for exposure to radiation, comprising administering to the mammal a composition comprising a therapeutically effective amount of a composition comprising an inducer of NF- κ B. The composition comprising an inducer of NF- κ B may be administered in combination with one or more radioprotectants. The one or more radioprotectants may be any agent that treats the effects of radiation exposure including, but not limited to, antioxidants, free radical scavengers and cytokines.

Inducers of NF- κ B may inhibit radiation-induced programmed cell death in response to damage in DNA and other cellular structures; however, inducers of NF- κ B may not deal with damage at the cellular level and may not prevent mutations. Free radicals and reactive oxygen species (ROS) are the major cause of mutations and other intracellular damage.

10 Antioxidants and free radical scavengers are effective at preventing damage by free radicals. The combination of an inducer of NF- κ B and an antioxidant or free radical scavenger may result in less extensive injury, higher survival, and improved health for exposure. Antioxidants and free radical scavengers that may be used in the practice of the invention include, but are not limited to, thiols, such as cysteine, cysteamine, glutathione and bilirubin; amifostine (WR-2721); vitamin A; vitamin C; vitamin E; and flavonoids such as orientin and vicenin derived from Indian holy basil (*Ocimum sanctum*).

15 Inducers of NF- κ B may also be administered in combination with a number of cytokines and growth factors that confer radioprotection by replenishing and/or protecting the radiosensitive stem cell populations. Radioprotection with minimal side effects may be achieved by the use of stem cell factor (SCF, c-kit ligand), Flt-3 ligand, and interleukin-1 fragment IL-1b-rd. Protection may be achieved through induction of proliferation of stem cells (all mentioned cytokines), and prevention of their apoptosis (SCF). The treatment allows 20 accumulation of leukocytes and their precursors prior to irradiation thus enabling quicker reconstitution of the immune system after irradiation. SCF efficiently rescues lethally irradiated mice with DMF in the range of 1.3-1.35 and is also effective against gastrointestinal syndrome. Flt-3 ligand also 25 provides strong protection in mice (70-80% 30-day survival at LD100/30, equivalent to DMF>1.2) and rabbits.

30 In addition, combinations of cytokines may provide enhanced radioprotection, such as: TPO combined with interleukin 4 (IL-4) and/or interleukin 11 (IL-11); GM-CSF combined with IL-3; G-CSF combined with Flt-3 ligand; 4F combination: SCF, Flt-3 ligand, TPO and IL-3; and 5F combination: 4F with addition of SDF-1.

35 In addition, gastrointestinal radioprotectors may be used, including transforming growth factor beta3 (TGF β 3), interleukin 11 (IL-11), and mentioned keratinocyte growth factor (KGF). While these radioprotectors also protect the intestine, they are likely to synergize with flagellin or flagellin related polypeptides since the results below show that flagellin and flagellin related polypeptides protect endothelium, while 40 these gastrointestinal radioprotectors protect epithelium of GI tract.

45 Several factors, while not cytokines by nature, stimulate the proliferation of immunocytes and may be used in combination with inducers of NF- κ B. For example, 5-AED (5-androstanediol) is a steroid that stimulates the expression of cytokines and increases resistance to bacterial and viral infections. A subcutaneous injection of 5-AED in mice 24 h before

irradiation improved survival with DMF=1.26. Synthetic compounds, such as ammonium tri-chloro(dioxoethylene-O, O⁻)-tellurate (AS-101), may also be used to induce secretion of numerous cytokines and for combination with inducers of NF-κB. Additional radioprotectors include, growth hormone (GH), thrombopoietin (TPO), interleukin 3 (IL-3), granulocyte-macrophage colony-stimulating factor (GM-CSF), granulocyte colony-stimulating factor (G-CSF), and stromal derived factor-1 (SDF-1).

Growth factors and cytokines may also be used to provide protection against gastrointestinal syndrome. Keratinocyte growth factor (KGF) promotes proliferation and differentiation in the intestinal mucosa, and increases the post-irradiation cell survival in the intestinal crypts. Hematopoietic cytokine and radioprotectant SCF may also increase intestinal stem cell survival and associated short-term organism survival.

Inducers of NF-κB may offer protection against both gastrointestinal (GI) and hematopoietic syndromes. Since mice exposed to 15 Gy of whole-body lethal irradiation die mostly from GI syndrome, a composition comprising an inducer of NF-κB and one or more inhibitors of GI syndrome may be more effective. Inhibitors of GI syndrome that may be used in the practice of the invention include, but are not limited to, cytokines such as SCF and KGF.

The composition comprising an inducer of NF-κB may be administered at any point prior to exposure to radiation including, but not limited to, about 48 hr, 46 hr, 44 hr, 42 hr, 40 hr, 38 hr, 36 hr, 34 hr, 32 hr, 30 hr, 28 hr, 26 hr, 24 hr, 22 hr, 20 hr, 18 hr, 16 hr, 14 hr, 12 hr, 10 hr, 8 hr, 6 hr, 4 hr, 3 hr, 2 hr, or 1 hr prior to exposure. The composition comprising an inducer of NF-κB may be administered at any point after exposure to radiation including, but not limited to, about 1 hr, 2 hr, 3 hr, 4 hr, 6 hr, 8 hr, 10 hr, 12 hr, 14 hr, 16 hr, 18 hr, 20 hr, 22 hr, 24 hr, 26 hr, 28 hr, 30 hr, 32 hr, 34 hr, 36 hr, 38 hr, 40 hr, 42 hr, 44 hr, 46 hr, or 48 hr after exposure to radiation.

f. Sepsis

This invention also relates to a method of preventing sepsis in a mammal comprising administering to the mammal a composition comprising a therapeutically effective amount of an agent that induces NF-κB activity. The agent that induces NF-κB activity may be administered in combination with other treatments.

Viral or bacterial infections may stimulate the innate immune system through Toll-like receptor (TLR) ligands. Macrophages may be protected and/or stimulated by flagellin and flagellin related polypeptides due to the presence of TLR5 on their surface. For example, a crucial step in the development of an anthrax infection is death of macrophages killed from within by *B. anthracis*. Protection of intestinal endothelium against various stresses using flagellin and flagellin related polypeptides may prevent GI cell death and also may prevent penetration of the GI wall by infectious agent, thereby preventing GI bleeding caused by infections such as Ebola. Other hemorrhagic viral infections may also be prevented by rescue of endothelium and gastrointestinal epithelium.

3. Agent

This invention also relates to an agent that induces NF-κB activity. The agent may be an artificially synthesized compound or a naturally occurring compound. The agent may be a low molecular weight compound, polypeptide or peptide, or a fragment, analog, homolog, variant or derivative thereof.

The agent may also be an NF-κB inducing cytokine including, but not limited to, IL2, IL6, TNF and TGFβ. The agent

may also be a prostaglandin. The agent may also be a growth factor including, but not limited to, KGF and PDGF. The agent may also be an antibody that induces NF-κB activity.

a. Flagellin

In one embodiment, the agent that induces NF-κB activity is flagellin. As shown in the Examples below, flagellin and flagellin related polypeptides possess strong pro-survival activity at the cellular level and for the organism as a whole. Interestingly, flagellin also stimulates natural killer (NK) cells and T-lymphocytes, which are the major components of anti-tumor immunity (Tsujimoto H, et. al., *J Leukoc Biol.* 2005 October; 78(4):888-97; Caron G., et. al., *J Immunol.* 2005 Aug. 1; 175(3):1551-7; Honko A N & Mizel S B, *Immunol Res.* 2005; 33(1):83-101). As a result, flagellin may be used as a radioprotectant in cancer treatments.

The present invention is also related to flagellin related polypeptides, such as those polypeptides described herein. As used herein, the term "flagellin" is intended to mean a flagellin or flagellin-related polypeptide from any source, including a variety of Gram-positive and Gram-negative bacterial species. The amino acid sequences of flagellin from 23 bacterial species are depicted in FIG. 7 of U.S. Patent Publication No. 2003/0044429, the contents of which are incorporated herein by reference. The nucleotide sequences encoding the flagellin polypeptides listed in FIG. 7 of U.S. 2003/0044429 are publicly available at sources including the NCBI Genbank database.

Flagellin is the major component of bacterial flagellum. Flagellin is composed of three domains (FIG. 9). Domain 1 (D1) and domain 2 (D2) are discontinuous and are formed when residues in the amino terminus and carboxy terminus are juxtaposed by the formation of a hairpin structure. The amino and carboxy terminus comprising the D1 and D2 domains is most conserved, whereas the middle hypervariable domain (D3) is highly variable. Studies with a recombinant protein containing the amino D1 and D2 and carboxyl D1 and D2 separated by an *Escherichia coli* hinge (ND1-2/ECH/CD2) indicate that D1 and D2 are bioactive when coupled to an ECH element. This chimera, but not the hinge alone, induced I_KB_α degradation, NF-κB activation, and NO and IL-8 production in two intestinal epithelial cell lines. The non-conserved D3 domain is on the surface of the flagellar filament and contains the major antigenic epitopes. The potent proinflammatory activity of flagellin may reside in the highly conserved N and C D1 and D2 regions.

Flagellin induces NF-κB activity by binding to Toll-like receptor 5 (TLR5). The TLR family is composed of at least 10 members and is essential in innate immune defense against pathogens. The innate immune system recognizes pathogen-associated molecular patterns (PAMPs) that are conserved on microbial pathogens. TLR may recognize a conserved structure that is particular to bacterial flagellin. The conserved structure may be comprised of a large group of residues that are somewhat permissive to variation in amino acid content. Smith et al., *Nat Immunol.* 4:1247-53 (2003) have identified 13 conserved amino acids in flagellin that are part of the conserved structure recognized by TLR5. The 13 conserved amino acids of flagellin important for TLR5 activity are shown in FIG. 24.

In a preferred embodiment, the flagellin is from a species of *Salmonella*, a representative example of which is *S. dublin* (encoded by GenBank Accession Number M84972) (SEQ ID NO: 1). In another preferred embodiment, the flagellin related-polypeptide is a fragment, variant, analog, homolog, or derivative of SEQ ID NO: 1, or combination thereof, that binds to TLR5 and induces TLR5-mediated activity, such as activation of NF-κB activity. A fragment, variant, analog,

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homolog, or derivative of flagellin may be obtained by rational-based design based on the domain structure of Flagellin and the conserved structure recognized by TLR5.

In a more preferred embodiment, the fragment, variant, analog, homolog, or derivative of SEQ ID NO: 1, or combination thereof, comprises at least 10, 11, 12, or 13 of the 13 conserved amino acids shown in FIG. 24 (positions 89, 90, 91, 95, 98, 101, 115, 422, 423, 426, 431, 436 and 452). In another more preferred embodiment, the amino- and carboxy-terminus of the fragment, variant, analog, homolog, or derivative of SEQ ID NO: 1, or combination thereof, is at least 30%, 35%, 40%, 45%, 50%, 55%, 60%, 65%, 70%, 75%, 80%, 85%, 90%, 95%, 97%, 98%, or 99% identical to amino acids 1-174 and 418-505 of SEQ ID NO: 1. FIG. 26 lists the percentage identity of the amino- and carboxy-terminus of flagellin with known TLR-5 stimulating activity, as compared to SEQ ID NO: 1.

Flagellin homologs may be a flagellin polypeptide from any Gram-positive or Gram-negative bacterial species including, but not limited to, the flagellin polypeptides disclosed in U.S. Pat. Pub. 2003/000044429, the contents of which are incorporated herein, and the flagellin peptides corresponding to the Accession numbers listed in the BLAST results shown in FIGS. 25A-D. Also contemplated, are fragments, variants, analogs and derivatives of flagellin homologs.

Flagellin fragments may be portions of a flagellin polypeptide that stimulate TLR5 activity. Numerous deletional mutants of flagellin have been made that retain at least some TLR5 stimulating activity. In addition to the deletional mutants disclosed in the Examples herein, representative deletional mutants include translation of GenBank Accession number D13689 missing amino acids 185-306 or 444-492, and translation of GenBank Accession number M84973 missing amino acids 179-415. Also contemplated, are homologs, variants, analogs and derivatives of flagellin fragments.

Flagellin variants include flagellin polypeptides with transposon insertions and changes to the variable D3 domain. The D3 domain may be substituted in part, or in whole, with a hinge or linker polypeptide that allows the D1 and D2 domains to properly fold such that the variant stimulates TLR5 activity. Representative examples of variant hinge elements may be found in the *E. coli* MukB protein and SEQ ID NOS: 3 and 4. Also contemplated, are fragments, homologs, analogs and derivatives of flagellin variants.

4. Composition

This invention also relates to a composition comprising a therapeutically effective amount of an inducer of NF- κ B. The composition may be a pharmaceutical composition, which may be produced using methods well known in the art. As described above, the composition comprising an inducer of NF- κ B may be administered to a mammal for the treatment of conditions associated with apoptosis including, but not limited to, exposure to radiation, side effect from cancer treatments, stress and cell aging. The composition may also comprise additional agents including, but not limited to, a radioprotectant or a chemotherapeutic drug.

a. Administration

Compositions of this invention may be administered in any manner including, but not limited to, orally, parenterally, sublingually, transdermally, rectally, transmucosally, topically, via inhalation, via buccal administration, intrapleurally, or combinations thereof. Parenteral administration includes, but is not limited to, intravenous, intraarterial, intraperitoneal, subcutaneous, intramuscular, intrathecal, and intraarticular. Transmucosally administration includes, but is not limited to

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intranasal. For veterinary use, the composition may be administered as a suitably acceptable formulation in accordance with normal veterinary practice. The veterinarian can readily determine the dosing regimen and route of administration that is most appropriate for a particular animal.

The composition may be administered prior to, after or simultaneously with a stress that triggers apoptosis, or a combination thereof. The composition may be administered from about 1 hour to about 48 hours prior to or after exposure to a stress that triggers apoptosis.

b. Formulation

Compositions of this invention may be in the form of tablets or lozenges formulated in a conventional manner. For example, tablets and capsules for oral administration may contain conventional excipients including, but not limited to, binding agents, fillers, lubricants, disintegrants and wetting agents. Binding agents include, but are not limited to, syrup, accacia, gelatin, sorbitol, tragacanth, mucilage of starch and polyvinylpyrrolidone. Fillers include, but are not limited to, lactose, sugar, microcrystalline cellulose, maizestarch, calcium phosphate, and sorbitol. Lubricants include, but are not limited to, magnesium stearate, stearic acid, talc, polyethylene glycol, and silica. Disintegrants include, but are not limited to, potato starch and sodium starch glycollate. Wetting agents include, but are not limited to, sodium lauryl sulfate). Tablets may be coated according to methods well known in the art.

Compositions of this invention may also be liquid formulations including, but not limited to, aqueous or oily suspensions, solutions, emulsions, syrups, and elixirs. The compositions may also be formulated as a dry product for constitution with water or other suitable vehicle before use. Such liquid preparations may contain additives including, but not limited to, suspending agents, emulsifying agents, non-aqueous vehicles and preservatives. Suspending agent include, but are not limited to, sorbitol syrup, methyl cellulose, glucose/sugar syrup, gelatin, hydroxyethylcellulose, carboxymethyl cellulose, aluminum stearate gel, and hydrogenated edible fats. Emulsifying agents include, but are not limited to, lecithin, sorbitan monooleate, and acacia. Non-aqueous vehicles include, but are not limited to, edible oils, almond oil, fractionated coconut oil, oily esters, propylene glycol, and ethyl alcohol. Preservatives include, but are not limited to, methyl or propyl p-hydroxybenzoate and sorbic acid.

Compositions of this invention may also be formulated as suppositories, which may contain suppository bases including, but not limited to, cocoa butter or glycerides. Compositions of this invention may also be formulated for inhalation, which may be in a form including, but not limited to, a solution, suspension, or emulsion that may be administered as a dry powder or in the form of an aerosol using a propellant, such as dichlorodifluoromethane or trichlorofluoromethane. Compositions of this invention may also be formulated transdermal formulations comprising aqueous or nonaqueous vehicles including, but not limited to, creams, ointments, lotions, pastes, medicated plaster, patch, or membrane.

Compositions of this invention may also be formulated for parenteral administration including, but not limited to, by injection or continuous infusion. Formulations for injection may be in the form of suspensions, solutions, or emulsions in oily or aqueous vehicles, and may contain formulation agents including, but not limited to, suspending, stabilizing, and dispersing agents. The composition may also be provided in a powder form for reconstitution with a suitable vehicle including, but not limited to, sterile, pyrogen-free water.

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Compositions of this invention may also be formulated as a depot preparation, which may be administered by implantation or by intramuscular injection. The compositions may be formulated with suitable polymeric or hydrophobic materials (as an emulsion in an acceptable oil, for example), ion exchange resins, or as sparingly soluble derivatives (as a sparingly soluble salt, for example).

c. Dosage

A therapeutically effective amount of the agent required for use in therapy varies with the nature of the condition being treated, the length of time that induction of NF- κ B activity is desired, and the age and the condition of the patient, and is ultimately determined by the attendant physician. In general, however, doses employed for adult human treatment typically are in the range of 0.001 mg/kg to about 200 mg/kg per day. The dose may be about 1 μ g/kg to about 100 μ g/kg per day. The desired dose may be conveniently administered in a single dose, or as multiple doses administered at appropriate intervals, for example as two, three, four or more subdoses per day. Multiple doses often are desired, or required, because NF- κ B activity in normal cells may be decreased once the agent is no longer administered.

The dosage of an inducer of NF- κ B may be at any dosage including, but not limited to, about 1 μ g/kg, 25 μ g/kg, 50 μ g/kg, 75 μ g/kg, 100 μ g/kg, 125 μ g/kg, 150 μ g/kg, 175 μ g/kg, 200 μ g/kg, 225 μ g/kg, 250 μ g/kg, 275 μ g/kg, 300 μ g/kg, 325 μ g/kg, 350 μ g/kg, 375 μ g/kg, 400 μ g/kg, 425 μ g/kg, 450 μ g/kg, 475 μ g/kg, 500 μ g/kg, 525 μ g/kg, 550 μ g/kg, 575 μ g/kg, 600 μ g/kg, 625 μ g/kg, 650 μ g/kg, 675 μ g/kg, 700 μ g/kg, 725 μ g/kg, 750 μ g/kg, 775 μ g/kg, 800 μ g/kg, 825 μ g/kg, 850 μ g/kg, 875 μ g/kg, 900 μ g/kg, 925 μ g/kg, 950 μ g/kg, 975 μ g/kg or 1 mg/kg.

This invention has multiple aspects, illustrated by the following non-limiting examples.

EXAMPLE 1

P53 Deficiency Accelerated Development of GI Syndrome in Mice

The primary cause of death from ionizing radiation (IR) of mammals depends on the radiation dose. At doses of up to 9-10 Gy, mice die 12-20 days later, primarily from lethal bone marrow depletion-hematopoietic (HP) syndrome. At this dose, irradiated mice can be rescued from lethality by bone marrow transplantation. Animals that receive >15 Gy die between 7-12 days after treatment (before hematopoietic syndrome can kill them) from complications of damage to the small intestine-gastrointestinal (GI) syndrome. In both cases of HP and GI syndromes, lethal damage of tissues starts from massive p53-dependent apoptosis. This observation allowed us earlier to suggest that p53 could be a determinant of radiation-induced death. Consistently, p53-deficient mice were resistant to doses of radiation that kill through HP syndrome, and lethality of wild type animals receiving 6-11 Gy of gamma radiation could be reduced by temporary pharmacological inhibition of p53 by the small molecule p53 inhibitor pifithrin-alpha (PFT) (Komarov et al 1999). Identification of p53 as a factor sensitizing tissues to genotoxic stress was further strengthened by demonstrating the p53 dependence of hair loss (alopecia) occurring as a result of experimental chemotherapy or radiation. Hence, based on previous observations, one would expect that p53 continues to play an important role in development of lethal GI syndrome after higher doses of IR. Surprisingly, p53-deficiency sensitizes mice to higher doses of IR causing lethal gastro-intestinal syndrome (FIG. 1). Continuous cell proliferation in the crypts

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of p53-deficient epithelium after IR correlates with accelerated death of damaged cells of crypt and rapid destruction of villi. p53 prolongs survival by inducing growth arrest in the crypts of small intestine thereby preserving integrity of the guts (FIG. 2). Thus, proapoptotic function of p53 promotes hematopoietic syndrome while its growth arrest function delays development of gastro-intestinal syndrome.

The dynamics of cell population in the small intestine have been analyzed in great detail. Cell proliferation in the epithelia of the gut is limited to the crypts where stem cells and early proliferating progenitors are located. After a couple of cell divisions, already differentiated descendants of crypt stem cells move up the villi to be shed at the villar tip. In the small intestine of the mouse, the entire "trip" of the cell (the proliferative compartment to the tip of the villus) normally takes between 3 and 5 days. Although reaction of the small intestine to gamma radiation has been well examined at a pathomorphological level, it still remains unclear what is the exact cause of GI lethality, including the primary event. Death may occur as a direct consequence of the damage of epithelial crypt cells and followed denudation of villi leading to fluid and electrolyte imbalance, bacteremia and endotoxemia. Besides inflammation and stromal responses, endothelial dysfunctions seem to be the important factors contributing to lethality. In summary, pharmacological suppression of p53 that was shown to be so effective as a method of protection from IR-induced HP syndrome, is useless (if not detrimental) against GI syndrome. Therefore, it is necessary to develop alternative approaches to radioprotection of epithelium of small intestine that will rely on another mechanism, such as, for example, activation of NF- κ B and subsequent inhibition of cell death.

EXAMPLE 2

Flagellin Delays Mouse Death Caused by IR-Induced GI Syndrome

Whole body irradiation of mice with 15 Gy gamma radiation caused death within 8 days from GI syndrome providing a conventional model of radiation induced damage of GI tract. To test whether flagellin was capable of protecting GI epithelium from IR, we tested the effect of i.v.-injected flagellin on the dynamics of mouse lethality after 15 Gy of radiation. We used a range of flagellin doses, all of which were significantly lower than the highest tolerable dose known from literature (300 μ g/mouse). Irradiation was done 4 hours post treatment. The results of a representative experiment are shown in FIG. 4. As expected, control irradiated mice (that received PBS i.v.) died between 5 and 8 days post-treatment. Animals that received flagellin lived significantly longer; the extension of animal survival correlated with the dose of flagellin. Pathomorphological analysis of the small intestine on day 7 after irradiation revealed dramatic differences between flagellin-treated and control groups (FIG. 5). Intravenous, intraperitoneal and subcutaneous delivery of 0.2 mg/kg of flagellin followed by 13 Gy irradiation afforded similar degree of protection, leading to 85-90% 30-day survival of mice (data not shown). While not being bound by theory, flagellin may be a radioprotectant due to its activation of NF- κ B, which presumably acts as an inhibitor of apoptotic death.

EXAMPLE 3

Flagellin Rescues Mice from Lethal IR-Induced Hematopoietic Syndrome

We next tested whether flagellin had an effect on mouse IR-induced death from HP syndrome that was experimentally

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induced by lower radiation doses (usually up to 11 Gy) that are incapable of causing lethal GI toxicity. The experiments were done similarly to the above-described ones (FIGS. 14 and 15), however, instead of 15 Gy, mice received 10 Gy, the dose that caused 100% killing in control group by day 13 (FIG. 6). Flagellin-treated group (5 µg/mouse) showed complete protection from this dose of IR surprisingly indicating that flagellin-mediated radioprotection acts not only against GI but also against HP IR-induced syndromes.

EXAMPLE 4

Time Dependence on the Protective Effect of Flagellin

Mice were next administered flagellin at different times prior to 13 Gy of gamma irradiation. The results of one of such experiments is shown in FIG. 7. The obtained results show that flagellin is effective as a radioprotectant from 13 Gy if injected 1-4 h before treatment.

In order to further estimate the dependence of radioprotective activity of flagellin on the time of treatment, mice were injected at several time points relative to the moment of gamma-irradiation. Experiments were performed essentially as explained above, using intraperitoneal injection of 5 µg/mouse (0.2 mg/kg) of full-length flagellin or, for control mice, 5 µg/mouse (0.2 mg/kg) of bacterial RNA polymerase. The experiments were performed using the NIH-Swiss mouse strain. The results show that flagellin provides ~90% survival after 13 Gy irradiation if injected at 1 or 2 hours before treatment (FIG. 7). Only -1 h graph is shown for clarity, however, both timepoints (-1 and -2 h) provide similar degree and dynamics of survival. The 4 h timepoint shows somewhat lower protection. Flagellin injected 24 hours before irradiation had no protective effect against 13 Gy induced death.

Interestingly, administration of flagellin 24 hours before 10 Gy gamma-irradiation provided 100% protection. While 13 Gy irradiation in mice primarily induces death from GI syndrome, 10 Gy-induced death is mostly mediated by hematopoietic syndrome. Accordingly, such long-term protection from 10 Gy irradiation may be mediated by enhanced proliferation or survival of hematopoietic stem cell induced by flagellin and/or long-living secondary cytokines.

EXAMPLE 5

Determination of LD_{50/30}, LD_{50/7} and DMF for Flagellin

We next obtained an estimate of radiation dose-dependent protection for flagellin. As shown above (FIG. 7), treatment with flagellin was sufficient for 100% protection against 10 Gy gamma-irradiation (this dose causes death from hematopoietic syndrome) and 90% 30-day survival at 13 Gy (both hematopoietic and GI syndromes). Experiments were performed as described above, using flagellin 5 µg/mouse (0.2 mg/kg), intraperitoneally injected 1 h before irradiation.

At 15 Gy, however, 100% 7-day survival was followed by delayed death after 13 days (0% 30-day survival), while control group had fully succumbed to GI syndrome by day 7 (FIG. 8). The kinetics of the flagellin-treated group mortality after 15 Gy irradiation is reminiscent of such of control group at 10 Gy, hinting at death caused by hematopoietic syndrome. The results provide an estimate of flagellin LD_{50/30} around

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13.5-14 Gy and DMF₃₀ of about 1.75-1.8. This degree of radioprotection is significantly higher than any reported for a natural compound.

EXAMPLE 6

Rational Design and Cloning of Flagellin Fragments

Salmonella flagellin, encoded by the FliC gene (SEQ ID NO: 2), is a strong activator of pro-survival NF-κB pathway. This is the most likely mechanism of its radioprotective action. Previous studies have shown that binding of flagellin to Toll-like receptor 5 (TLR5) on the cell surface is a necessary step that triggers activation of NF-κB. The domain structure of *Salmonella* flagellin is described in sufficient detail in the literature (FIG. 9). Moreover, previous structural studies of flagellin-TLR5 complex (FIG. 10) provide the ability to distinguish between domains that are essential or dispensable for binding and thus NF-κB activation. Protein minimization may provide reduced immune response after repeated administration of flagellin-related polypeptides. This may be achieved, in part, due to lower immunogenicity of low molecular weight proteins and smaller number of immunogenic epitopes available.

The domains needed for TLR5 binding may be located exclusively in the evolutionary conserved N- and C-terminal domains of bacterial flagellins. The hypervariable domain (amino acids 178-402) does not come into close contact with TLR5. As was demonstrated previously, replacement of this domain with a flexible linker peptide did not disrupt binding to TLR5. In addition, N-terminal and C-terminal coiled-coil polymerization domains (amino acids 1-55, 456-505) do not bind to TLR5 and likely are dispensable (see modified N and C termini B and B', respectively, as shown herein). Also, another fragment N-terminus lacking all domains but major N-terminal α-helix that actually binds TLR5 (amino acids 56-100) may be sufficient for binding.

Accordingly, three types of N-termini (A, B, C) and two types of C-termini (A', B'), connected with a flexible linker (SEQ ID NOS: 3 and 4) taken from pGEX-KG cloning vector (SEQ ID NOS: 5 and 6) were combined into expression constructs to produce several possible flagellin fragments (Table 1). In addition, constructs representing separate N-termini (A, B, C) and glutathione-S-transferase (GST)-fusions of C-termini (GST-A', GST-B') were prepared. All constructs were cloned in the pRSETb bacterial expression vector and 6xHis-tagged proteins were produced and purified for further experiments (FIG. 11).

TABLE 1

Name	Structure	DNA	Protein
AA'	(1-177)-Linker-(402-505)	SEQ ID NO: 7	SEQ ID NO: 8
AB'	(1-177)-Linker-(402-450)	SEQ ID NO: 9	SEQ ID NO: 10
BA'	(56-177)-Linker-(402-505)	SEQ ID NO: 11	SEQ ID NO: 12
BB'	(56-177)-Linker-(402-450)	SEQ ID NO: 13	SEQ ID NO: 14
CA'	(56-100)-Linker-(402-505)	SEQ ID NO: 15	SEQ ID NO: 16
CB'	(56-100)-Linker-(402-450)	SEQ ID NO: 17	SEQ ID NO: 18
A	(1-177)	SEQ ID NO: 19	SEQ ID NO: 20
B	(56-177)	SEQ ID NO: 21	SEQ ID NO: 22
C	(56-100)	SEQ ID NO: 23	SEQ ID NO: 24
GST-A'	GST-Linker-(402-505)	SEQ ID NO: 25	SEQ ID NO: 26
GST-B'	GST-Linker-(402-450)	SEQ ID NO: 27	SEQ ID NO: 28

EXAMPLE 7

Selection Of Biologically Active Flagellin Fragments

Since the radioprotective activities of flagellin appear to be NF-κB dependent, we tested the ability of the flagellin frag-

ments to induce NF- κ B translocation to the nucleus and binding to its target sites in DNA. This was tested by electrophoretic mobility shift assay (EMSA) using nuclear extracts from flagellin- and fragment-treated A549 lung cancer cells and labeled synthetic NF- κ B binding kB oligonucleotide.

Only flagellin itself and fragments AA', AB', and BA' were capable of inducing NF- κ B translocation (FIG. 12). The level of translocation is comparable for flagellin and fragments AA', AB', and BA'. The hypervariable domain does not appear to be necessary for NF- κ B translocation, while the presence of at least one polymerization domain, N- or C-terminal, is required. Mixtures of the N- and C-terminal fragments (A+A', A+B') were inactive.

While translocation of NF- κ B to the nucleus is a crucial step in induction of NF- κ B-regulated inhibitors of apoptosis, it is not sufficient in itself. To directly test the ability of selected fragments to induce expression of NF- κ B-regulated genes, we performed reporter assay experiments. Flagellin and the AA', BB', A' and B' fragments were used for treatment of H116 human colon cancer cells carrying luciferase gene under a NF- κ B-responsive promoter. The reporter construct contained three NF- κ B-binding sites from the E-selectin promoter combined with a Hsp70 minimal promoter that is routinely used for the detection of NF- κ B status of cells. Luciferase activity was measured in cell lysates six hours after addition of flagellin or its truncated fragments into the medium. TNF was used as positive control. The results of a representative experiment are shown in FIG. 13 and indicates that flagellin and fragment AA' are capable of NF- κ B activation, whereas fragments BB', GST-A' and GST-B' are not.

EXAMPLE 8

Further Optimization of Flagellin Fragments

We further minimized the AA' flagellin fragment by producing additional fragments through stepwise removal of peptide fragments from its N-terminal half (Table 2). Electrophoretic mobility shift assays were performed as described above using nuclear extracts from flagellin- and fragment-treated HT29 human colon cancer cells and labeled synthetic NF- κ B binding kB oligonucleotide. NF- κ B binding activity in HT29 cells was stimulated with TNFa (10 ng/ml), or flagellin fragments (1 mg/ml) for 15 min. As shown in FIG. 14, fragments AA'n1-170, AA'n54-170, AA'n1-163 and AA'n54-163 each induce NF- κ B translocation, with levels comparable to that of flagellin for AA'n1-170, AA'n54-170 and AA'n1-163.

TABLE 2

Name	Structure	DNA	Protein
AA' n54-177	(54-177)-Linker-(402-505)	SEQ ID NO: 11	SEQ ID NO: 12
AA' n1-170	(1-170)-Linker-(402-505)	SEQ ID NO: 29	SEQ ID NO: 30
AA' n54-170	(54-170)-Linker-(402-505)	SEQ ID NO: 31	SEQ ID NO: 32
AA' n1-163	(1-163)-Linker-(402-505)	SEQ ID NO: 33	SEQ ID NO: 34
AA' n54-163	(54-163)-Linker-(402-505)	SEQ ID NO: 35	SEQ ID NO: 36

In order to study the ability of the AA' fragments to directly activate NF- κ B-regulated transcription, we performed reporter assay experiments as described above for a wide range of concentrations of flagellin, original AA' and AA'-derived fragments. As shown above, AA' and AA'n1-170

induce NF- κ B-regulated transcription at the level comparable to such of flagellin over the studied range of concentrations (FIG. 15, left). AA' and AA'n1-170 are more active than flagellin in the very low concentration range (FIG. 15, right), possibly due to their reduced molecular weight. The results with fragment AA'n1-170 show that AA'-derived flagellin fragments may be made with a portion of the N-terminal domain removed without significant loss of activity and may be used as effective radioprotectors.

The above experiments (EMSA and reporter activation assay) were repeated with flagellin and AA' fragments subjected to 30 minutes boiling and renaturation before being applied to cells. The results were comparable to those obtained without boiling (data not shown). This shows that the observed differences in flagellin fragment activity may not be caused by changes in protein stability.

EXAMPLE 9

In Vivo Comparison of Radioprotective Properties of Flagellin and Flagellin Fragments

As shown above, full-length flagellin provides protection from both hematopoietic and gastrointestinal syndromes. The radioprotective potential of flagellin fragments was similarly tested after gamma-irradiation with 11 Gy (dose that induces hematopoietic syndrome-associated mortality in mice) or 14 Gy (dose that causes death from GI syndrome). Mice (10 animals per group) were injected subcutaneously with 5.0 μ g/mouse (0.2 mg/kg) of flagellin or its fragments, AA' or BB', and gamma-irradiated 1 hour later.

The degree of radioprotection displayed by the AA' fragment is at least comparable to full-length flagellin (FIG. 16). Both the AA' fragment and full-length flagellin showed 100% 30-day survival for mice irradiated with 11 Gy and 14 Gy. Meanwhile, 0% of mice injected with the BB' fragment survived to 30 days. This is expected since the BB' fragment is incapable of inducing NF- κ B in vitro. These results show that significant reduction in the size of flagellin (about 40% removed) may be achieved without a decrease in the degree of radioprotection. In addition, the ability to predict radioprotective potential from results of in vitro NF- κ B activation is confirmed.

EXAMPLE 10

Identification of Cellular Targets of Flagellin-Mediated Radioprotection

Tissue samples of intestinal mucosa were taken 5 days after 14 Gy irradiation from mice pretreated with flagellin and control mice. Control animals were treated with 5.0 μ g/mouse (0.2 mg/kg) bacterial RNA-polymerase. Pathomorphological analysis of the small intestine reveals reduction of the size of crypts and villi and a number of the cells with condensed apoptotic nuclei in control mouse and near-normal morphology in the treated mouse (data not shown). Tissue samples (small intestine and skin from the back) were also obtained from mice treated with the AA' fragment. The results shown in FIG. 17 are areas of typical morphology observed over a set of at least 3 mice. After treatment with flagellin and the AA' fragment, mice demonstrated near-normal intestinal morphology with preservation of the villi/crypt structure (FIG. 17A).

In addition to purely histological observation of cell death and survival, we performed more specialized tests of apoptotic cell death in intestinal tissue using a TUNEL assay,

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which detects apoptosis-associated DNA fragmentation. These experiments allowed us to define with a high degree of probability cellular populations that are depleted by radiation and rescued by flagellin fragment treatment. The earliest radiation-induced alterations detectable in the small intestine after treatment with IR is apoptosis occurring in vascular endothelial cells of villi, which is seen as early as 5 hours post treatment (FIG. 17B). This apoptosis, which is believed to be critical for radiosensitivity of the small intestine, was almost completely blocked in the mice pretreated with the AA' fragment (FIG. 17B, bottom panel). Degeneration of villi and crypts, occurring within the next several days post treatment and greatly suppressed in AA' fragment-treated animals, comes as a consequence of injury of blood vessels. Effective protection of endothelial cells of the small intestine by the flagellin fragment may be due to expression of TLR5 in these cells.

Remarkably, the AA' fragment and flagellin also prevented the radiation-induced disappearance of sebaceous glands located at the base of skin hair follicles (FIG. 27C). These results further confirm the suitability of AA' fragment for radioprotection and for the prevention of radiation-induced hair loss.

EXAMPLE 11

Protection from Supralethal Radiation

In order to explore the limits of radioprotection provided by the AA' fragment, we irradiated mice with 17 Gy and 20 Gy single doses of total body gamma-radiation. The experiment was performed as described above using inactive flagellin fragment (CB) as a negative control.

As expected, we observed a 100% mortality in both groups at 17 and 20 Gy (FIG. 18). However, death was significantly delayed in both cases by administration of the AA' fragment. Most remarkably, the kinetics of death at 17 Gy in control mice conform to GI syndrome (6-7 day mortality), while death of mice treated with the AA' fragment appear to be mediated by hematopoietic syndrome (10-15 day mortality). This shows that flagellin and flagellin fragments may protect against the GI syndrome at doses as high as 17 Gy. In addition, this shows that even further radioprotection may be obtained by flagellin and flagellin fragments combined with hematopoietic radioprotectors.

EXAMPLE 12

Immunogenicity and Repetitive Administration Studies

Overall immunogenicity of a protein may determines its suitability for repeated use. Antibodies generated by immune system are capable of reducing the therapeutic activity of the protein and also may induce anaphylactic reaction upon second exposure if IgE antibodies are produced against the protein. Thus, any reduction in the amount and variety of antibodies compared to full-length flagellin is an improvement. Accordingly, after repeated introduction of flagellin or its fragments we monitored; a) efficiency of radioprotection afforded at second exposure; b) local and general allergic reactions; and c) antibody titer.

We tested the ability of AA' to protect mice that were exposed to it. A group of 20 NIH-Swiss mice were subcutaneously injected with 5 µg/mouse (0.2 mg/kg) of AA'. A second injection of an equal dose of AA' was administered after 21 (10 mice) and 28 days (another 10 mice), with the time

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elapsed being sufficient for formation of antibodies. The second injection of AA' was followed by 13 Gy of whole-body gamma irradiation (1 h post-injection). 100% 30-day survival was observed in both groups, as it was observed with mice that had no previous exposure to AA' (data not shown). These results show that activity of AA' is not diminished over long-term repeated administration and reaffirm its potential for multiple-use applications. Also, no local allergic reaction or anaphylaxis was observed either with flagellin or AA'.

We also performed an ELISA determination of antibody titers in order to quantify the effect that AA' has on the immune status of the organism. 96-well plates were coated with flagellin or AA', 20 mg/ml, 50 ml/well, and incubated overnight at +4°C. Blood serum samples collected from mice were added to the wells in several dilutions and incubated overnight followed by 6 hrs reaction with secondary goat anti-mouse IgG HPO-conjugate antibodies. Measurements were performed using a spectrophotometer with a 414 nm filter. The antibody titers determined for individual mice and average titers are shown in FIG. 19 and FIG. 20.

AA' induces far lower antibody levels in mice (FIG. 19), on the order of 0.8 mg/ml serum at 21 day and about 10% more at 28 days. Flagellin, on another hand (FIG. 20), induces a high titer of antibodies, around 20 mg/ml, at both 21 and 28 days. Overall, this shows that removal of the hypervariable domain sharply reduces the immunogenicity of AA' compared to the original protein (approximately 25x). FIG. 19 also shows that the majority of AA'-specific antibodies are capable of recognizing flagellin. This confirms that the rational design of AA' does not produce a sizable number of new immunogenic epitopes while removing >95% of the immunogenicity of the original protein.

EXAMPLE 13

Acute Toxicity Studies of Flagellin and AA'

The lethal dose of *Salmonella* flagellin is between 1 mg/kg (systemic inflammation) and 10 mg/kg (100% lethality). We subcutaneously administered increasing doses of AA' to mice (4 mice per dose group) at 0.5, 1, 2, 4 and 8 mg/kg. Due to the lower (~60%) molecular weight, 8 mg/kg of AA' correspond to a molar-equivalent dose of 13.3 mg/kg of flagellin. Several days after administration at all doses, no visible detrimental effects were observed, such as mortality, morbidity or signs of systemic inflammation such as reduced activity and fever. This shows that the pro-inflammatory effect of AA' is negligible compared to full-length flagellin, especially considering that AA' provides an efficient radioprotection at 0.2 mg/kg. The reduced toxicity may be due to the absence of the central pro-inflammatory domain in the AA' fragment.

EXAMPLE 14

Protection from Fractioned Irradiation by AA'

Repetitive irradiation within a short period of time may be common, for example, in space radiation events and in clinical radiotherapy regimens. We tested the ability of the AA' flagellin fragment to protect mice from sub-lethal (4 treatments of 3 Gy) and 100% lethal (4 treatments of 4 Gy) regimens of fractionated gamma-irradiation. Fragment AA' or saline buffer was given to NIH-Swiss female mice before every irradiation (once a day for 4 days). AA' was administered as described above for single-dose irradiation (5 µg/mouse, given subcutaneously 1 h before irradiation).

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The results in FIG. 21 show that AA' provides significant protection against repetitive doses of radiation received within a short timeframe. The cumulative dose of fractionated radiation that is still compatible with 100% 30-day survival after AA' treatment is comparable to such obtained in single-dose irradiation scenarios.

EXAMPLE 15

AA' Protects Normal Tissues without Compromising the Anti-Tumor Therapeutic Effect of Radiation

The ultimate test of a potential radioprotective agent in cancer treatment is tumor selectivity, its ability to protect normal tissues while providing no or little protection to the tumor. We injected 10 NIH-Swiss mice subcutaneously, in both flanks (20 tumors total), with 2×10^6 cells of syngeneic sarcoma cell line model (NIH3T3-derived and spontaneously transformed sarcoma with p53 inactivated by dominant negative inhibitor GSE56). When tumors reached the size of 5-7 mm in diameter (day 5), the mice were injected subcutaneously with 0.2 mg/kg of AA' or saline vehicle and irradiated 1 hour later with 4 Gy of total-body γ -irradiation (3×4.3 Gy=12.9 Gy total dose). Injections and irradiations were done at days 5, 6 and 7.

As the results show in FIG. 11, AA' enhanced the radiation-induced shrinkage of tumors. By day 18, all the irradiated tumor-bearing mice died from acute radiation toxicity whereas 100% of mice that obtained both radiotherapy and AA' were both cured and survived the treatment. Similar result was obtained with another syngeneic tumor model—B16 melanoma cells (FIG. 11, right panel). Surprisingly, even in unirradiated mice, AA' administration caused a decrease in the growth rate of tumors. This may be due to AA'-induced immunostimulating, which is known to be caused by other ligands of Toll-like receptors. These results indicate that AA' increases the tolerance of mice to radiation with no effect on the radiosensitivity of two types of tumors, thus opening the possibility of combining radiotherapy with AA' to improve treatment outcome.

EXAMPLE 16

Radioprotective Mechanisms of AA' and LPS are Different

Lipopolysaccharide of gram-negative bacteria (LPS) is a ligand of another Toll-like receptor, TLR4. LPS is a strong inducer of NF- κ B and a subsequent cascade of cytokines. LPS is known as a radioprotective compound, but its high toxicity makes its use unfeasible (radioprotective dose is very close to the lethal dose). One of the major mechanisms underlying the radioprotection by LPS is the activation of cyclooxygenase 2 (COX-2) that, in turn, drives the synthesis of GI-protective prostaglandins. The possibility that radio-protection by TLR5 also relies on COX-2 activity was tested by administering s.c. LPS (2 mg/kg), AA' (0.2 mg/kg) or vehicle 1 hr before irradiation of NIH-Swiss mice in combination with i.p. injection of 1 mg/kg of NS398, a synthetic COX-2 inhibitor, or the corresponding vehicle. The mice were then treated with 13 Gy of total-body γ -irradiation. NS398 completely abolished LPS-mediated radioprotection but not the radioprotection of AA' (FIG. 12). This result shows that AA' does not significantly rely on COX-2 for its activity and induces radioprotection by a mechanism different from the mechanism of LPS-mediated protection.

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EXAMPLE 17

AA' Protects Multiple Mouse Strains

We have extensively confirmed that AA' protects NIH-Swiss and ICR mice from radiation. To confirm that the radioprotection activity of AA' is not confined to a few mouse strains, several additional strains of mice with dissimilar origins were tested for protection by AA': 129/Sv, DBA/2 (relatively radioresistant), Balb/c (relatively radiosensitive) and Balb/cxDBA/2 F1 hybrid CD2F1. Experimental groups were injected with 0.2 mg/kg AA' 30 minutes before irradiation, while control groups were injected with vehicle (PBS).

All groups of mice (8-10 mice each, 8-12 week old females) were exposed to 10 Gy of single-dose, whole-body gamma-irradiation. Survival of mice at days 10 and 30 is shown. The results are shown in FIG. 27 as a cone graph. At 10 days, only Balb/c mice display mortality, which is drastically reduced by AA' administration (0% vs. 100% survival). At day 30, all tested strains display improved survival (0-25% vs. 50-100%) after AA' administration.

EXAMPLE 18

Pharmacokinetics of AA'

Pharmacokinetic parameters (effective concentration and the duration of drug the presence in the organism) may be important for route, dose and time of drug administration. The pharmacokinetics of CBLB502 (AA') were thus tested for four common routes of injection: intravenous (i.v.), subcutaneous (s.c.), intraperitoneal (i.p.) or intramuscular (i.m.). A radioprotective dose of CBLB502 (AA'), 0.2 mg/kg, was injected in 12-15 week old ICR mice and plasma samples were collected at the specified times after injection (at least 3 mice/point). The levels of CBLB502 in plasma were measured by sandwich ELISA using known concentrations of CBLB502 spiked in the control ICR plasma for calibration. The results are shown in FIG. 28 and FIG. 29.

The results show that the highest levels and longest persistence of CBLB502 in plasma are provided by intramuscular or intraperitoneal injection. After intramuscular injection, significant (>5 ng/ml) levels of CBLB502 are observed in mouse plasma >3 hours. Intravenous injection leads to a more rapid disappearance of CBLB502 from the bloodstream.

EXAMPLE 19

Influence of AA' on Gamma-Irradiation Induced Cell Death and Growth Inhibition in A549 Cells

The A549 human lung adenocarcinoma cell line is reported to respond to flagellin by activation of NF- κ B DNA-binding activity (Tallant T., et. al., *BMC Microbiol.* 2004 Aug. 23; 4:33). We decided to check whether this activation translates in the protection of cells from γ -IR in cell growth inhibition assay.

Tumor cells were seeded in wells of three 96-well plates in 3 different densities (0.5×10^4 , 1×10^4 and 2×10^4 cells/well, producing single-cell, spare or semi-contact layer). After cells had attached to plastic, CBLB502 (2 μ g/ml) was added to the wells of non-irradiated cells, or 15 min prior to 7 Gy or 10 Gy of gamma-irradiation. Control wells received equal volume of vehicle (PBS). All points were done in quadruplicate. 72 hours after irradiation, medium was replaced with methylene blue in 50%-methanol and the relative numbers of viable cells in wells were measured using spectrophotometer

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at 650 nm. The results are shown in FIG. 30. This experiment was also repeated with fixed dose of 1×10^4 A549 cells/well, CBLB502 was added 1 hour before 5, 10 or 15 Gy of gamma-irradiation (data not shown). We observed a similar effect of flagellin in all tested experiment conditions.

Gamma-irradiation induced a dose-dependent reduction in the number of A549 cells plated at all three densities (up to 60% as compared to non-irradiated control wells). CBLB502 had no or slight effect on cell numbers, with or without gamma-irradiation. This indicates that tumor cells are not significantly protected by CBLB502 (AA') from radiation. This effect may be due to tumor cells having constitutively active NF- κ B pathway or some other mechanism.

EXAMPLE 20**Influence of AA' on Gamma-Irradiation Induced Cell Death and Growth Inhibition in Multiple Cell Lines**

Based on the results using A549 cells, several additional tumor cell lines (human melanoma Mel-7 and Mel-29, colon cancer HCT116, lung cancer HT1080), immortalized kidney epithelial cells (NKE) and normal mouse aortal endothelial cells (MAEC) were tested in growth inhibition assay after 10 and 15 Gy of gamma-irradiation, as compared with intact control, with or without pretreatment with CBLB502. Cells were seeded in 96-well plates night before the treatments. CBLB502 (2 μ g/ml) was added to the wells 4 hrs, 1 hr or 10 min before irradiation (all points were done in quadruplicate). 48 hours later, methylene blue staining was performed to determine the relative amount of the viable cells in the wells. All three time-points had shown the same effect (results for CBLB502 added 1 hr before irradiation are shown in FIG. 31). The percent of growth inhibition was calculated from the OD650 in control non-irradiated wells, taken as 0% inhibition.

Both human melanoma cell lines and MAEC cells were rather resistant to gamma-irradiation and showed only slight (<20%) growth inhibition after both 10 and 15 Gy comparing with intact cells (0 Gy). NKE, HT1080 and HCT116 cells showed up to 40% of growth inhibition after gamma-irradiation. Remarkably, CBLB502 had no or only a slight inhibitory effect on tumor cell growth, irradiated or not. The experiment was repeated twice. In addition, similar results were obtained on tested lung adenocarcinoma H1299 and prostate cancer CWR22 (data not shown). This indicates that there is no significant protection provided by CBLB502 to the tumor cell lines against radiation-induced cell death.

EXAMPLE 21**Influence of Irradiation and AA' on BrdU Incorporation in Small Intestinal Crypts**

Besides a direct inhibition of apoptosis, temporary halt of proliferation followed by repair may be an alternative mechanism of radioprotection, and has been described for other radioprotectors such as TGF- β 3 (Booth D., et al., *Int J Cancer*. 2000 Apr. 1; 86(1):53-9). Accordingly, we decided to examine the possible influence of CBLB502 (AA') on the proliferative activity of the cells in small intestine (with and without irradiation) during the first hours after its administration (FIG. 32). CBLB502 or PBS was injected i.p. in mice, followed after 30 min by 15 Gy irradiation (if used). 2 hr after injection (1.5 hr after irradiation if it was applied), BrdU was injected intraperitoneally. Samples of small intestine were obtained after additional 1.5 hours.

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Without irradiation, BrdU was incorporated at high levels in the nuclei of cells in the intestinal crypts of untreated NIH-Swiss mice (FIG. 32, top left), whereas DNA synthesis (as measured by BrdU incorporation) was nearly undetectable in the crypts of CBLB502-treated mice (FIG. 32, top right). In vehicle-treated irradiated mice, the incorporation of BrdU was lower than in control mice. Importantly, the level of BrdU incorporation was strongly reduced by CBLB502, possibly indicating quick (S phase) growth arrest, as opposed to later (G2 phase) irradiation-induced growth arrest. Therefore, cytostatic activity of CBLB502 or flagellin may be an additional mechanism of radioprotection of small intestine.

EXAMPLE 22**Duration of AA'-Mediated Growth Arrest and Reduced BrdU Incorporation**

We next determined the duration of the CBLB502-induced growth arrest in small intestine. CBLB502 or PBS was injected i.p. in mice, BrdU was injected 1 or 4 hours later and samples of small intestine were obtained after additional 1.5 hours from several mice (samples from three mice are shown) (FIG. 33).

Incorporation of BrdU in intestine was reduced as compared to control if BrdU was injected after 1 hour (as it was shown in the previous experiment where BrdU was injected after 2 hr). NIH-Swiss, ICR and Balb/c mice displayed a similar degree of CBLB502-mediated block of BrdU incorporation (Balb/c samples are shown in FIG. 33). If BrdU was injected 4 hr after injection of CBLB502, the levels of incorporation/DNA synthesis were even higher than in control. This indicates that inhibition of intestinal stem cell proliferation by CBLB502 may be temporary and may be quickly resolved (by 4 hours), followed by a period of increased proliferation (possibly due to the partial synchronization of cells).

EXAMPLE 23**Influence of AA' on BrdU Incorporation in Colonic Crypts**

The colon is much less radiosensitive than small intestine. To further examine the relationship between reduced proliferation in the small intestine and radioprotection, we determined the effect of CBLB502 on BrdU incorporation in the colon. CBLB502 or PBS was injected i.p. in mice, BrdU was injected 1 hour later and samples of small intestine were obtained after additional 1.5 hours (FIG. 34).

Unlike in the small intestine, CBLB502 has no effect on BrdU incorporation in colon. This is surprising since TLR5 is plentiful in both organs. The difference in effects may be due to the higher amount of symbiotic bacteria in the colon, which may mask the effect of additional TLR5 signaling induced by CBLB502.

EXAMPLE 24**Comparison of Radioprotective Potential by Route of Administration**

We next tested radioprotection provided by FliC flagellin administered via several routes: intravenous (i.v.), intraperitoneal (i.p.), intramuscular (i.m.), subcutaneous (s.c.) and gavage. For parenteral (non-gavage) routes, mice were injected with 0.2 mg/kg of FliC flagellin dissolved in PBS or

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vehicle, followed 1 hr later by 13 Gy irradiation. In gavage delivery experiment, 5 mice were given to swallow an increased dose (50 µg) of FliC in 50 µl of PBS 1 h before 13 Gy gamma-irradiation. Both experiments were done in 8-10 week old female NIH-Swiss mice, 5-10 mice/group.

All tested routes besides gavage afforded similar degree of protection, leading to 85-90% 30-day survival of mice (data not shown). No protection against radiation was provided by gavage delivery, which may be due to digestion of the protein by the gastrointestinal environment. In addition, flagellin receptor, TLR5, is absent on the luminal side of intestinal epithelium that is exposed to intestinal contents (Gewirtz A T., et. al., *J Immunol.* 2001 Aug. 15; 167(4):1882-5)

EXAMPLE 25

Effect of AA' on the Morphology of Small Intestine

Flagellin (and CBLB502) may induce NF-κB activity via binding to TLR5. Accordingly, CBLB502-mediated radio-

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with or without CBLB502. This indicates that the presence of TLR5 may be necessary for CBLB502-mediated radioprotection in the small intestine.

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EXAMPLE 26

Flagellin Derivatives

Additional flagellin variants were produced based on the domain structure shown in FIG. 36. The flagellin variants were then tested along with some of the variants discussed above for NF-κB stimulating activity (Table 3). A549 cells were left unstimulated or stimulated with TNF (10 ng/ml) as indicated or 1 µg/ml of purified flagellin or the various indicated flagellin derivatives for 45 min and whole cell extracts prepared as described in Example 7. EMSA assays were preformed and the NF-κB DNA-protein complex detected as described in Example 7.

TABLE 3

Name	N-terminal	C-terminal	DNA	Protein	NF-κB Stimulation
AA'	1-176	402-505	SEQ ID NO: 7	SEQ ID NO: 8	Yes
AB'	1-176	402-450	SEQ ID NO: 9	SEQ ID NO: 10	Yes
BA'	54-176	402-505	SEQ ID NO: 11	SEQ ID NO: 12	Yes
BB'	54-176	402-450	SEQ ID NO: 13	SEQ ID NO: 14	No
CA'	54-100	402-505	SEQ ID NO: 15	SEQ ID NO: 16	No
CB'	54-100	402-450	SEQ ID NO: 17	SEQ ID NO: 18	No
AA'n1-170	1-170	402-505	SEQ ID NO: 29	SEQ ID NO: 30	Yes
AA'n54-170	54-170	402-505	SEQ ID NO: 31	SEQ ID NO: 32	Yes
AB'n1-170	1-170	402-450	SEQ ID NO: 37	SEQ ID NO: 38	Yes
AA'n1-163	1-163	402-505	SEQ ID NO: 33	SEQ ID NO: 34	Yes
AA'n54-163	54-163	402-505	SEQ ID NO: 35	SEQ ID NO: 36	Yes
AB'n1-163	1-163	402-450	SEQ ID NO: 39	SEQ ID NO: 40	Yes
AA'n1-129	1-129	402-505	SEQ ID NO: 41	SEQ ID NO: 42	Yes
AA'n54-129	54-129	402-505	SEQ ID NO: 43	SEQ ID NO: 44	Yes
AB'n1-129	1-129	402-450	SEQ ID NO: 45	SEQ ID NO: 46	untested
AB'n54-129	54-129	402-450	SEQ ID NO: 47	SEQ ID NO: 48	Untested
AA'n1-100	1-100	402-505	SEQ ID NO: 49	SEQ ID NO: 50	untested
AB'n1-100	1-100	402-450	SEQ ID NO: 51	SEQ ID NO: 52	untested
AA'n1-70	1-70	402-505	SEQ ID NO: 53	SEQ ID NO: 54	No
AB'n1-70	1-70	402-450	SEQ ID NO: 55	SEQ ID NO: 56	No
A	1-176		SEQ ID NO: 19	SEQ ID NO: 20	No
B	54-176		SEQ ID NO: 21	SEQ ID NO: 22	No
C	54-100		SEQ ID NO: 23	SEQ ID NO: 24	No
GST-A'		402-505	SEQ ID NO: 25	SEQ ID NO: 26	No
GST-B'		402-450	SEQ ID NO: 27	SEQ ID NO: 28	No

protection may be dependent on the presence and activity of TLR5. MOLF/Ei mice are a known natural model of TLR5 deficiency (Sebastiani G., et. al., *Genomics.* 2000 Mar. 15; 64(3):230-40). To verify that CBLB502-mediated radioprotection is indeed TLR5-dependent, we tested the protection of small intestine from radiation by CBLB502 in MOLF/Ei and NIH-Swiss mice (FIG. 35). Both strains of mice were given 0.2 mg/kg CBLB502 (AA') or PBS 0.5 hr before 15 Gy of gamma-irradiation. The samples of small intestine were obtained 4 days after irradiation, stained by hematoxylin-eosin and subjected to pathomorphological analysis.

In NIH-Swiss (TLR5 wild type) mice, CBLB502 pretreatment led to preservation of intestinal morphology (long villae, normal crypts) as compared to short villae and disappearance of normal crypt structure in PBS-treated mice. Meanwhile, in TLR5-deficient MOLF/Ei mice the administration of CBLB502 had no improving effect on intestinal morphology after 15 Gy of gamma-irradiation: short villae and destruction of the normal crypt structure was observed,

The results in Table 3 indicate that flagellin variants with at least one polymerization domain (aa1-50 or aa 450-505) that linked to domains contained within the amino-terminal region (aa1-176) and those of the carboxy terminus (aa 402-505) are capable of stimulating NF-κB and would thus be expected to be radioprotectors. Physical linkage of the recognition domains may be required for activity as domains supplied unlinked in trans fail to activate NF-κB. As an alternative to the linking of the domains in a single polypeptide, the domains may be linked using a linker, which is a molecule that is used to join two molecules. The linker may be capable of forming covalent bonds or high-affinity non-covalent bonds to both molecules. Suitable linkers are well known to those of ordinary skill in the art and include, but are not limited to, straight or branched-chain carbon linkers, heterocyclic carbon linkers, or peptide linkers. The linkers may be joined to the constituent amino acids through their side groups (e.g., through a disulfide linkage to cysteine).

The region between amino acids 163 and 176 may be required for activity when the carboxyl polymerization

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domain (aa 450-505) is absent. Since this region is dispensable for activity when the carboxyl polymerization domain is present it may be involved in stabilizing the derivative. The region between amino acids 70 and 129 may be important for activation and may be involved in derivative recognition. The region between amino acids 402 and 450 may also be required

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for activity. The domains identified above are located within three large α -helices (located within amino acids 54-129 and 402-450) and, to produce an active derivative, may need to form a ring-like structure (with or without polymerization domain).

SEQUENCE LISTING

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<211> LENGTH: 505
<212> TYPE: PRT
<213> ORGANISM: Salmonella dublin

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Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg Leu
 20          25          30

Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
 35          40          45

Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala
 50          55          60

Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly
 65          70          75          80

Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu Ser
 85          90          95

Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser Ile
100         105         110

Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser Asn
115         120         125

Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln Met
130         135         140

Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp Leu
145         150         155         160

Gln Lys Ile Asp Val Lys Ser Leu Gly Leu Asp Gly Phe Asn Val Asn
165         170         175

Gly Pro Lys Glu Ala Thr Val Gly Asp Leu Lys Ser Ser Phe Lys Asn
180         185         190

Val Thr Gly Tyr Asp Thr Tyr Ala Ala Gly Ala Asp Lys Tyr Arg Val
195         200         205

Asp Ile Asn Ser Gly Ala Val Val Thr Asp Ala Ala Ala Pro Asp Lys
210         215         220

Val Tyr Val Asn Ala Ala Asn Gly Gln Leu Thr Thr Asp Asp Ala Glu
225         230         235         240

Asn Asn Thr Ala Val Asp Leu Phe Lys Thr Thr Lys Ser Thr Ala Gly
245         250         255

Thr Ala Glu Ala Lys Ala Ile Ala Gly Ala Ile Lys Gly Gly Lys Glu
260         265         270

Gly Asp Thr Phe Asp Tyr Lys Gly Val Thr Phe Thr Ile Asp Thr Lys
275         280         285

Thr Gly Asp Asp Gly Asn Gly Lys Val Ser Thr Thr Ile Asn Gly Glu
290         295         300

Lys Val Thr Leu Thr Val Ala Asp Ile Ala Thr Gly Ala Ala Asp Val
305         310         315         320

Asn Ala Ala Thr Leu Gln Ser Ser Lys Asn Val Tyr Thr Ser Val Val

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37**38**

-continued

325	330	335
Asn Gly Gln Phe Thr Phe Asp Asp Lys Thr Lys Asn Glu Ser Ala Lys		
340	345	350
Leu Ser Asp Leu Glu Ala Asn Asn Ala Val Lys Gly Glu Ser Lys Ile		
355	360	365
Thr Val Asn Gly Ala Glu Tyr Thr Ala Asn Ala Thr Gly Asp Lys Ile		
370	375	380
Thr Leu Ala Gly Lys Thr Met Phe Ile Asp Lys Thr Ala Ser Gly Val		
385	390	395
Ser Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys Ser Thr Ala		
405	410	415
Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val		
420	425	430
Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr		
435	440	445
Asn Leu Gly Asn Thr Val Thr Asn Leu Asn Ser Ala Arg Ser Arg Ile		
450	455	460
Glu Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Lys Ala Gln		
465	470	475
Ile Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val		
485	490	495
Pro Gln Asn Val Leu Ser Leu Leu Arg		
500	505	

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<211> LENGTH: 1518

<212> TYPE: DNA

<213> ORGANISM: *Salmonella dublin*

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ttcagtcct cactgagttc cgcttatttagcg cgtctgtcct ctggctgcgt tatcaacagc	120
gcgaaagacg atgcggcagg ccaggcgatt gctaaccgcg tcaacttctaa tatcaaaggc	180
ctgactcagg cttcccgtaa cgctaaccgac ggcatttcta ttgcgcagac cactgaaggt	240
gcgctgaatg aaatcaacaa caacctgcag cgtgtgegtg agttgtctgt tcaggccact	300
aacgggacta actctgattc cgatctgaaa tctatccagg atgaaattca gcaacgtctg	360
gaagaaaatcg atcgcgttcc taatcagact caatattaacg gtgttaaagt cctctctcag	420
gacaaccaga tgaaaatcca gggttgtct aacgatggtg aaaccattac catcgatctg	480
caaaaaatttgc atgtgaaaag ctttggcctt gatgggttca atgttaatgg gccaaaagaa	540
gcgacagtgg gtgatctgaa atccagcttc aagaatgtta cgggttacga cacctatgca	600
gcgggtgccg ataaatatcg tgttagatatt aattccgggtc ctgttagtgac ttagcgcgca	660
gcaccggata aagtatatgt aaatgcagca aacgggtcagt taacaactga ccatgcggaa	720
aataaacactg cgggtgatct ctttaagacc actaaatcta ctgctggtac cgctgaagcc	780
aaagcgatag ctgggtccat taaaggttgtt aaggaaggagg ataccttgc ttataaaggc	840
gtgactttta ctattgatac aaaaacttgtt gatgacggta atggtaaggt ttctactacc	900
atcaatgggtt aaaaagttac gttaactgtc gctgatatttgc ccactggcgc ggcggatgtt	960
aatgctgcta ctttacaatc aagcaaaaat gtttatacat ctgttagtcaa cggtcagttt	1020
acttttgatg ataaaaccaa aaacgagagt gcgaaactttt ctgatttgaa agcaaacaat	1080

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gctgttaagg	gcgaaagtaa	aattacagta	aatggggctg	aatatactgc	taacgccacg	1140
ggtgataaga	tcacccttagc	tggcaaaacc	atgtttattt	ataaaaacagc	ttctggcgta	1200
agtacattaa	tcaatgaaga	cgcgtccgca	gccaagaaaa	gtaccgctaa	cccactggct	1260
tcaattgatt	ctgcattgtc	aaaagtggac	gcagttcggt	cttctctggg	ggcaattcaa	1320
aaccgttttg	attcagccat	taccaacatt	ggcaatacgg	taaccaatct	gaactcccgcg	1380
cgtageccgta	tcgaagatgc	tgactatgca	acggaagttt	ctaatatgtc	taaagcgcag	1440
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<212> TYPE: PRT
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic sequence (peptide linker)

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<212> TYPE: PRT
<213> ORGANISM: Artificial Sequence
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<223> OTHER INFORMATION: Synthetic sequence (peptide linker)

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<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic sequence (peptide linker)

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<210> SEQ ID NO 6
<211> LENGTH: 46
<212> TYPE: DNA
<213> ORGANISM: Artificial Sequence
<220> FEATURE:
<223> OTHER INFORMATION: Synthetic sequence (peptide linker)

<400> SEQUENCE: 6

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<211> LENGTH: 990
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<213> ORGANISM: Salmonella dublin

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aacagcctgt	cgctgttgac	ccagaataac	ctgaacaaat	ctcagtcctc	actgagttcc	180

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gctattggc gtctgtccctc tggctcgct atcaacagcg cgaaagacga tgccggcaggc 240
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gctaaccgacg gcattttctat tgcgccagacc actgaagggtg cgctgaatga aatcaacaac 360
aacctgcagc gtgtgcgtga gttgtctgtt caggccacta acgggactaa ctctgattcc 420
gatctgaaat ctatccagga tgaatttcag caacgtctgg aagaaatcga tcgcgtttct 480
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cttggcccttg atgggttcaa tggtaattcc cggggattt ccgggtggtgg tggtggatt 660
ctagactcca tgggtacatt aatcaatgaa gacgctgccc cagccaagaa aagtaccgt 720
aaccctactgg cttcaattga ttctgcattt tcaaaagtgg acgcagttcg ttcttctctg 780
ggggcaattt aaaaccgtt tgattcagcc attaccaacc ttggcaatac ggttaaccaat 840
ctgaactccg cgcgttagccg tategaagat gctgactatg caacggaaat ttctaatatg 900
tctaaagcgc agattctgca gcaggctgtt acttccgttc tggcgcaggc taaccagggtt 960
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<211> LENGTH: 329
<212> TYPE: PRT
<213> ORGANISM: *Salmonella dublin*

<400> SEQUENCE: 8

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20 25 30

Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln
35 40 45

Asn Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg
50 55 60

Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
65 70 75 80

Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
85 90 95

Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu
100 105 110

Gly Ala Leu Asn Glu Ile Asn Asn Asn Leu Gln Arg Val Arg Glu Leu
115 120 125

Ser Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser
130 135 140

Ile Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser
145 150 155 160

Asn Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln

Met Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp

Leu Gln Lys Ile Asp Val Lys Ser Leu Gly Leu Asp Gly Phe Asn Val

Asn Ser Pro Gly Ile Ser Gly Gly Gly Gly Gly Ile Leu Asp Ser Met

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Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys Ser Thr Ala
225 230 235 240

Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val
245 250 255

Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr
260 265 270

Asn Leu Gly Asn Thr Val Thr Asn Leu Asn Ser Ala Arg Ser Arg Ile
275 280 285

Glu Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Lys Ala Gln
290 295 300

Ile Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val
305 310 315 320

Pro Gln Asn Val Leu Ser Leu Leu Arg
325

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<212> TYPE: DNA
<213> ORGANISM: Salmonella dublin

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aacagectgt cgctgttgac ccagaataac ctgaacaaat ctcagtcctc actgagttcc        180
gttattgagc gtctgtccctc tggctctgcgt atcaacagcg cgaaagacga tgccggcaggc        240
caggcgattt ctaaccgctt cacttctaat atcaaaggcc tgactcaggc ttcccgtaac        300
gctaaccgacg gcatttctat tgccgacacc actgaagggtc cgctgaatga aatcaacaaac        360
aacctgcaggc gtgtgcgtga gttgtctgtt caggccacta acgggactaa ctctgattcc        420
gatctgaaat ctatccagga tgaaattcag caacgtctgg aagaaaatcga tcgcgttct        480
aatcagactc aatttaacgg tggtaaagtc ctctctcagg acaaccagat gaaaatcag        540
gttgggtcata acgatggtga aaccattacc atcgatctgc aaaaaattga tggaaaagc        600
cttggccttg atgggttcaa tggtaattcc cgggaattt cgggtggtgg tggtaatt        660
cttagactcca tgggtacatt aatcaatgaa gacgctgccg cagccaagaa aagtaccgct        720
aacccactgg cttcaattga ttctgcattt tcaaaagtgg acgcagttcg ttcttcttg        780
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<212> TYPE: PRT
<213> ORGANISM: Salmonella dublin

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Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Asp Lys Asp
20                    25                    30

Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln
35                    40                    45

Asn Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg
50                    55                    60

Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
65                    70                    75                    80
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Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
 85 90 95

Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu
 100 105 110

Gly Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu
 115 120 125

Ser Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser
 130 135 140

Ile Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser
 145 150 155 160

Asn Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln
 165 170 175

Met Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp
 180 185 190

Leu Gln Lys Ile Asp Val Lys Ser Leu Gly Leu Asp Gly Phe Asn Val
 195 200 205

Asn Ser Pro Gly Ile Ser Gly Gly Gly Ile Leu Asp Ser Met
 210 215 220

Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys Ser Thr Ala
 225 230 235 240

Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val
 245 250 255

Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr
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Asn Leu

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 <212> TYPE: DNA
 <213> ORGANISM: *Salmonella dublin*

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ctgactcagg	cttcccgtaa	cgctaacgcac	ggcatttctta	ttgcgcagac	cactgaaggt	180
gcgcgtgaatg	aaatcaacaa	caacctgcag	cgtgtgcgtg	agttgtctgt	tcaggccact	240
aacgggacta	actctgattc	cgtatctgaaa	tctatccagg	atgaaattca	gcaacgtctg	300
gaagaaaatcg	atcgcgtttc	taatcagact	caatttaacg	gtgttaaagt	cctctctcag	360
gacaaccaga	tgaaaatcca	ggttggtgct	aacgatggtg	aaaccattac	catcgatctg	420
caaaaaatgtt	atgtgaaaag	ccttggcatt	gatgggtca	atgttaatc	cccgaaaatt	480
tccgggttgt	gtgggtgaat	tcttagactcc	atgggtacat	taatcaatga	agacgctgcc	540
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gacgcagttc	gttcttctct	ggggcaattt	caaaaccgtt	ttgattcagc	cattaccaac	660
cttggcaata	cggttaacaa	tctgaactcc	gcgcgttagcc	gtatcgaaga	tgctgactat	720
gcaacggaag	tttctaatat	gtctaaagcg	cagattctgc	agcaggctgg	tacttccgtt	780
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 <212> TYPE: PRT

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20					25				30					

Pro	Phe	Thr	Ser	Asn	Ile	Lys	Gly	Leu	Thr	Gln	Ala	Ser	Arg	Asn	Ala
35					40				45						

Asn	Asp	Gly	Ile	Ser	Ile	Ala	Gln	Thr	Thr	Glu	Gly	Ala	Leu	Asn	Glu
50					55				60						

Ile	Asn	Asn	Asn	Leu	Gln	Arg	Val	Arg	Glu	Leu	Ser	Val	Gln	Ala	Thr
65					70			75			80				

Asn	Gly	Thr	Asn	Ser	Asp	Ser	Asp	Leu	Lys	Ser	Ile	Gln	Asp	Glu	Ile
85					90				95						

Gln	Gln	Arg	Leu	Glu	Glu	Ile	Asp	Arg	Val	Ser	Asn	Gln	Thr	Gln	Phe
100					105				110						

Asn	Gly	Val	Lys	Val	Leu	Ser	Gln	Asp	Asn	Gln	Met	Lys	Ile	Gln	Val
115					120				125						

Gly	Ala	Asn	Asp	Gly	Glu	Thr	Ile	Thr	Ile	Asp	Leu	Gln	Lys	Ile	Asp
130					135			140							

Val	Lys	Ser	Leu	Gly	Leu	Asp	Gly	Phe	Asn	Val	Asn	Ser	Pro	Gly	Ile
145					150			155			160				

Ser	Gly	Gly	Gly	Ile	Leu	Asp	Ser	Met	Gly	Thr	Leu	Ile	Asn		
165					170			175							

Glu	Asp	Ala	Ala	Ala	Lys	Lys	Ser	Thr	Ala	Asn	Pro	Leu	Ala	Ser	
180					185			190							

Ile	Asp	Ser	Ala	Leu	Ser	Lys	Val	Asp	Ala	Val	Arg	Ser	Ser	Leu	Gly
195					200			205							

Ala	Ile	Gln	Asn	Arg	Phe	Asp	Ser	Ala	Ile	Thr	Asn	Leu	Gly	Asn	Thr
210					215				220						

Val	Thr	Asn	Leu	Asn	Ser	Ala	Arg	Ser	Ile	Glu	Asp	Ala	Asp	Tyr	
225					230			235			240				

Ala	Thr	Glu	Val	Ser	Asn	Met	Ser	Lys	Ala	Gln	Ile	Leu	Gln	Gln	Ala
245					250			255							

Gly	Thr	Ser	Val	Leu	Ala	Gln	Ala	Asn	Gln	Val	Pro	Gln	Asn	Val	Leu
260					265			270							

Ser	Leu	Leu	Arg												
275															

<210> SEQ ID NO 13

<211> LENGTH: 666

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 13

atgcgggggtt ctcatcatca tcatcatcat ggtatggcta gcatgactgg tggacagcaa 60

atgggtcggg atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggc 120

ctgactcagg cttcccgtaa cgctaaccgac ggcatttcta ttgcgcagac cactgaaggt 180

gcgcgtaatg aaatcaacaa caaacctgcag cgtgtgcgtg agttgtctgt tcaggccact 240

aacggggacta actctgtatcc cgatctgaaa tctatccagg atgaaattca gcaacgtctg 300

gaagaaaatcg atcgcgttcc taatcagact caatattaacg gtgttaaagt cctctctcag 360

gacaaccaga tgaaaatcca gggtggtgct aacgatggtg aaaccattac catcgatctg 420

-continued

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caaaaaattg atgtgaaaag ccttggcctt gatgggttca atgttaattc cccggaaatt    480
tccgggttgtg gtgggtgaat tctagactcc atgggtacat taatcaatga agacgctgcc    540
gcagccaaga aaagtaccgc taacccactg gttcaattt attctgcatt gtcaaaaatg    600
gacgcagttc gttttctct gggggcaatt caaaaccgtt ttgattcagc cattaccaac    660
cttttag                                         666

```

<210> SEQ ID NO 14

<211> LENGTH: 221

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 14

```

Met Arg Gly Ser His His His His His Gly Met Ala Ser Met Thr
1           5           10          15

```

```

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp
20          25          30

```

```

Pro Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala Ser Arg Asn Ala
35          40          45

```

```

Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly Ala Leu Asn Glu
50          55          60

```

```

Ile Asn Asn Asn Leu Gln Arg Val Arg Glu Leu Ser Val Gln Ala Thr
65          70          75          80

```

```

Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser Ile Gln Asp Glu Ile
85          90          95

```

```

Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser Asn Gln Thr Gln Phe
100         105         110

```

```

Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln Met Lys Ile Gln Val
115         120         125

```

```

Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp Leu Gln Lys Ile Asp
130         135         140

```

```

Val Lys Ser Leu Gly Leu Asp Gly Phe Asn Val Asn Ser Pro Gly Ile
145         150         155         160

```

```

Ser Gly Gly Gly Ile Leu Asp Ser Met Gly Thr Leu Ile Asn
165         170         175

```

```

Glu Asp Ala Ala Ala Lys Lys Ser Thr Ala Asn Pro Leu Ala Ser
180         185         190

```

```

Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val Arg Ser Ser Leu Gly
195         200         205

```

```

Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn Leu
210         215         220

```

<210> SEQ ID NO 15

<211> LENGTH: 603

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 15

```

atgcgggtt ctcatcatca tcatcatcat ggtatggata gcatgactgg tggacagcaa    60
atgggtcggg atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggc    120
ctgactcagg cttcccgtaa cgctaacgac ggcatttcta ttgcgcagac cactgaaggt    180
gcgcgtgaatg aaatcaacaa caacctgcag cgtgtgcgtg agttgtctgt tcaggccact    240
tccccggaa tttccgggtgg tggtgggtgg attcttagact ccatgggtac attaatcaat    300

```

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gaagacgctg ccgcagccaa gaaaagtacc gctaaccac tggcttcaat tgattctgca	360
ttgtcaaaag tggacgcagt tcgttcttct ctgggggcaa ttcaaaacgg ttttgatca	420
gccattacca accttggcaa tacggtaacc aatctgaact ccgcgcgttag ccgtatcgaa	480
gtatgtact atgcaacgga agtttctaat atgtctaaag cgcatgttct gcagcaggct	540
gttacttccg ttctggcgca ggctaacccag gttccgcaaa acgtcctctc tttactgcgt	600
tag	603

<210> SEQ ID NO 16

<211> LENGTH: 200

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 16

Met Arg Gly Ser His His His His His Gly Met Ala Ser Met Thr			
1	5	10	15

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp			
20	25	30	

Pro Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala Ser Arg Asn Ala			
35	40	45	

Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly Ala Leu Asn Glu			
50	55	60	

Ile Asn Asn Asn Leu Gln Arg Val Arg Glu Leu Ser Val Gln Ala Thr			
65	70	75	80

Ser Pro Gly Ile Ser Gly Gly Gly Ile Leu Asp Ser Met Gly			
85	90	95	

Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys Ser Thr Ala Asn			
100	105	110	

Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val Arg			
115	120	125	

Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn			
130	135	140	

Leu Gly Asn Thr Val Thr Asn Leu Asn Ser Ala Arg Ser Arg Ile Glu			
145	150	155	160

Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Lys Ala Gln Ile			
165	170	175	

Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val Pro			
180	185	190	

Gln Asn Val Leu Ser Leu Leu Arg		
195	200	

<210> SEQ ID NO 17

<211> LENGTH: 438

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 17

atgcgggtt ctcatcatca tcatacatcat ggtatggcta gcatgactgg tggacagcaa	60
atgggtcggtt atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggc	120
ctgactcagg cttcccgtaa cgctaacgac ggcatttcta ttgcgcagac cactgaaggt	180
gcgcgtaat aaatcaacaa caacctgcag cgtgtgcgtg agttgtctgt tcaggccact	240
tccccggaa ttcccggtgg tggtggtgaa attctagact ccatgggtac attaatcaat	300
gaagacgctg ccgcagccaa gaaaagtacc gctaaccac tggcttcaat tgattctgca	360

-continued

ttgtcaaaag tggacgcagt tcgttcttct ctgggggcaa ttcaaaaccg ttttgattca 420

gccattatcca accttttag 438

<210> SEQ ID NO 18

<211> LENGTH: 145

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 18

Met	Arg	Gly	Ser	His	His	His	His	His	Gly	Met	Ala	Ser	Met	Thr
1														
														15

Gly	Gly	Gln	Gln	Met	Gly	Arg	Asp	Leu	Tyr	Asp	Asp	Asp	Lys	Asp
20														30

Pro	Phe	Thr	Ser	Asn	Ile	Lys	Gly	Leu	Thr	Gln	Ala	Ser	Arg	Asn	Ala
35															45

Asn	Asp	Gly	Ile	Ser	Ile	Ala	Gln	Thr	Thr	Glu	Gly	Ala	Leu	Asn	Glu
50															60

Ile	Asn	Asn	Asn	Leu	Gln	Arg	Val	Arg	Glu	Leu	Ser	Val	Gln	Ala	Thr
65															80

Ser	Pro	Gly	Ile	Ser	Gly	Gly	Gly	Ile	Leu	Asp	Ser	Met	Gly	
85														95

Thr	Leu	Ile	Asn	Glu	Asp	Ala	Ala	Ala	Lys	Lys	Ser	Thr	Ala	Asn
100														110

Pro	Leu	Ala	Ser	Ile	Asp	Ser	Ala	Leu	Ser	Lys	Val	Asp	Ala	Val	Arg
115															125

Ser	Ser	Leu	Gly	Ala	Ile	Gln	Asn	Arg	Phe	Asp	Ser	Ala	Ile	Thr	Asn
130															140

Leu

145

<210> SEQ ID NO 19

<211> LENGTH: 639

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 19

atgcggggtt ctcatcatca tcatacatcat ggtatggcta gcatgactgg tggacagcaa 60

atgggtcggg atctgtacga cgatgacgat aaggatccga tggcacaagt cattaataca 120

aacagcctgt cgctgttgac ccagaataac ctgaacaaat ctcagtcctc actgagttcc 180

gtatttgagc gtctgtcctc tggctctgcgt atcaacagcgc cgaaagacga tgcggcaggc 240

caggcgattg ctaaccgctt cacttctaat atcaaaggtc tgactcaggc ttcccgtaac 300

getaacgacg gcatttctat tgccgacacc actgaagggtg cgctgaatga aatcaacaac 360

aacctgcagc gtgtgcgtga gttgtctgtt caggccacta acgggactaa ctctgattcc 420

gatctgaat ctatccagga taaaattcag caacgtctgg aaaaaatcga tcgcgtttct 480

aatcagactc aattnaacgg tggtaaaggc ctgtctcagg acaaccagat gaaaatccag 540

gttgggtcata acgatggtga aaccattacc atcgatctgc aaaaaattga tgtgaaaagc 600

cttggccttg atgggttcaa tggtaattcc ccgggatga 639

<210> SEQ ID NO 20

<211> LENGTH: 212

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 20

-continued

Met Arg Gly Ser His His His His His His Gly Met Ala Ser Met Thr
 1 5 10 15
 Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp
 20 25 30
 Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln
 35 40 45
 Asn Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg
 50 55 60
 Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
 65 70 75 80
 Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
 85 90 95
 Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu
 100 105 110
 Gly Ala Leu Asn Glu Ile Asn Asn Asn Leu Gln Arg Val Arg Glu Leu
 115 120 125
 Ser Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser
 130 135 140
 Ile Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser
 145 150 155 160
 Asn Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln
 165 170 175
 Met Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp
 180 185 190
 Leu Gln Lys Ile Asp Val Lys Ser Leu Gly Leu Asp Gly Phe Asn Val
 195 200 205
 Asn Ser Pro Gly
 210

<210> SEQ ID NO 21
 <211> LENGTH: 480
 <212> TYPE: DNA
 <213> ORGANISM: *Salmonella dublin*
 <400> SEQUENCE: 21
 atgcgggggtt ctcatcatca tcatcatcat ggtatggcta gcatgactgg tggacagcaa 60
 atgggtcggg atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggt 120
 ctgactcagg cttcccgtaa cgctaaccac ggcatttctta ttgcgcagac cactgaaggt 180
 gcgcgtaatg aaatcaacaa caacctgcag cgtgtgcgtg agttgtctgt tcaggccact 240
 aacgggacta actctgattc cgatctgaaa tctatccagg atgaaattca gcaacgtctg 300
 gaagaaaatcg atcgcgttcc taatcagact caatttaacg gtgttaaagt cctgtctcag 360
 gacaaccaga tgaaaatcca gggttgtgct aacgatggtg aaaccattac catcgatctg 420
 caaaaaatcg atgtaaaaag ccttggcctt gatgggttca atgttaattc cccgggatga 480

<210> SEQ ID NO 22
 <211> LENGTH: 159
 <212> TYPE: PRT
 <213> ORGANISM: *Salmonella dublin*
 <400> SEQUENCE: 22
 Met Arg Gly Ser His His His His His Gly Met Ala Ser Met Thr
 1 5 10 15
 Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp

-continued

20 25 30

Pro Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala Ser Arg Asn Ala
 35 40 45

Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly Ala Leu Asn Glu
 50 55 60

Ile Asn Asn Asn Leu Gln Arg Val Arg Glu Leu Ser Val Gln Ala Thr
 65 70 75 80

Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser Ile Gln Asp Glu Ile
 85 90 95

Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser Asn Gln Thr Gln Phe
 100 105 110

Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln Met Lys Ile Gln Val
 115 120 125

Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp Leu Gln Lys Ile Asp
 130 135 140

Val Lys Ser Leu Gly Leu Asp Gly Phe Asn Val Asn Ser Pro Gly
 145 150 155

<210> SEQ ID NO 23

<211> LENGTH: 252

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 23

atgcgggggtt ctcatcatca tcatacatcat ggtatggcta gcatgactgg tggacagcaa 60

atggggcggg atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggt 120

ctgactcagg cttcccgtaa cgctaacgc acgcatttcta ttgcgcagac cactgaaggt 180

gcgcgtgaatg aaatcaacaa caaacctgcag cgtgtgcgtg agttgtctgt tcaggccact 240

tccccgggat ga 252

<210> SEQ ID NO 24

<211> LENGTH: 83

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 24

Met Arg Gly Ser His His His His His Gly Met Ala Ser Met Thr
 1 5 10 15

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Asp Lys Asp
 20 25 30

Pro Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala Ser Arg Asn Ala
 35 40 45

Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly Ala Leu Asn Glu
 50 55 60

Ile Asn Asn Asn Leu Gln Arg Val Arg Glu Leu Ser Val Gln Ala Thr
 65 70 75 80

Ser Pro Gly

<210> SEQ ID NO 25

<211> LENGTH: 1038

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 25

atgtccctta tactaggtta ttggaaaatt aagggccttg tgcaaccac tcgacttctt 60

-continued

ttgaaatatac ttgaagaaaa atatgaagag catttgtatg agcgcatga aggtgataaa	120
tggcgaaaca aaaagtttga attgggtttg gagtttccca atcttccta ttatattgat	180
ggtgatgtta aattaacaca gtctatggcc atcatacgat atatacgta caagcacaa	240
atgttgggtg gttgtccaaa agagcgtgca gagattcaa tgcttgaagg agcggtttg	300
gatattagat acgggtttc gagaatttgcata tagttaaag actttgaaac tctcaagtt	360
gatatttcta gcaagctacc tgaaatgtcg aaaatgttc aagatcgat atgtcataaa	420
acatatttaa atggtcatca tgtaacccat cctgacttca tgttgtatga cgcttgc	480
gttggtttat acatggaccc aatgtgcgtg gatgcgttcc caaaatttagt ttgtttaaa	540
aaacgtattt aagctatccc acaaatttgcata aagtacttgc aatccagcaa gtatatacg	600
tggccttgc agggctggca agccacgttt ggtggggcc accatcctcc aaaatcgat	660
ctgggtccgc gtggatcccc gggatttcc ggtgggggtg gtggattct agactccatg	720
ggtacattaa tcaatgaaga cgctgccgca gccaagaaaa gtaccgctaa cccactggct	780
tcaatttgcatttgc aaaaatggac gcagtgcgtt ctgcgttccc ggcaatttca	840
aaccgttttgc attcagccat taccaacctt ggcaatacgg taaccaatct gaactcccg	900
cgttagccgta tcgaagatgc tgactatgc acggaagttt ctaatatgtc taaagcgcag	960
attctgcagc aggctggta ttccgttctg ggcaggcttcc accagggttcc gcaaaacgtc	1020
ctctctttac tgcgttag	1038

<210> SEQ_ID NO 26

<211> LENGTH: 345

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 26

Met Ser Pro Ile Leu Gly Tyr Trp Lys Ile Lys Gly Leu Val Gln Pro			
1	5	10	15

Thr Arg Leu Leu Glu Tyr Leu Glu Glu Lys Tyr Glu Glu His Leu			
20	25	30	

Tyr Glu Arg Asp Glu Gly Asp Lys Trp Arg Asn Lys Lys Phe Glu Leu			
35	40	45	

Gly Leu Glu Phe Pro Asn Leu Pro Tyr Tyr Ile Asp Gly Asp Val Lys			
50	55	60	

Leu Thr Gln Ser Met Ala Ile Ile Arg Tyr Ile Ala Asp Lys His Asn			
65	70	75	80

Met Leu Gly Cys Pro Lys Glu Arg Ala Glu Ile Ser Met Leu Glu			
85	90	95	

Gly Ala Val Leu Asp Ile Arg Tyr Gly Val Ser Arg Ile Ala Tyr Ser			
100	105	110	

Lys Asp Phe Glu Thr Leu Lys Val Asp Phe Leu Ser Lys Leu Pro Glu			
115	120	125	

Met Leu Lys Met Phe Glu Asp Arg Leu Cys His Lys Thr Tyr Leu Asn			
130	135	140	

Gly Asp His Val Thr His Pro Asp Phe Met Leu Tyr Asp Ala Leu Asp			
145	150	155	160

Val Val Leu Tyr Met Asp Pro Met Cys Leu Asp Ala Phe Pro Lys Leu			
165	170	175	

Val Cys Phe Lys Lys Arg Ile Glu Ala Ile Pro Gln Ile Asp Lys Tyr			
180	185	190	

Lys Ser Ser Lys Tyr Ile Ala Trp Pro Leu Gln Gly Trp Gln Ala	
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195 200 205

Thr Phe Gly Gly Asp His Pro Pro Lys Ser Asp Leu Val Pro Arg
 210 215 220

Gly Ser Pro Gly Ile Ser Gly Gly Gly Ile Leu Asp Ser Met
 225 230 235 240

Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys Ser Thr Ala
 245 250 255

Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val
 260 265 270

Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr
 275 280 285

Asn Leu Gly Asn Thr Val Thr Asn Leu Asn Ser Ala Arg Ser Arg Ile
 290 295 300

Glu Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Lys Ala Gln
 305 310 315 320

Ile Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val
 325 330 335

Pro Gln Asn Val Leu Ser Leu Leu Arg
 340 345

<210> SEQ ID NO 27

<211> LENGTH: 873

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 27

atgtccctta tactaggtta ttggaaaatt aagggccttg tgcaaccac tcgacttctt	60
ttggaatatac ttgaagaaaa atatgaagag catttgtatg agcgcgatga aggtgataaaa	120
tggcgaaaca aaaagtttga attgggtttg gagttccca atcttcctta ttatattgtat	180
ggtgtatgtta aattaacaca gtctatggcc atcatacggtt atatacgctga caagcacaac	240
atgttgggtg gttgtccaaa agagcgtgca gagatttcaa tgcttgaagg agcggttttg	300
gatatttagat acgggttttc gagaatttgcata tatacgaaac tctcaaagtt	360
gattttctta gcaagctacc tgaaatgctg aaaatgttgc aagatcggtt atgtcataaaa	420
acatatttaa atgggtatca tgtaacccat cctgacttca tgttgtatga cgctcttgc	480
gttgttttat acatggaccc aatgtgcctg gatgcgttcc caaaattagt ttgttttaaa	540
aaacgtatttgc aagctatccc acaaattgtat aagtacttgc aatccagcaa gtatatacgca	600
tggccttgc agggctggca agccacgttt ggtggggcg accatcctcc aaaatcgat	660
ctgggtccgc gtggatcccc gggatttcc ggtgggtggtg gtggattctt agactccatg	720
ggtacattaa tcaatgaaga cgctgccca gccaagaaaa gtaccgctaa cccactggct	780
tcaatttgcatttgc aaaaatggac gcagttcggtt cttctctggg ggcaattcaa	840
aaccgttttg attcagccat taccaacctt tag	873

<210> SEQ ID NO 28

<211> LENGTH: 290

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 28

Met Ser Pro Ile Leu Gly Tyr Trp Lys Ile Lys Gly Leu Val Gln Pro	
1 5 10 15	

Thr Arg Leu Leu Leu Glu Tyr Leu Glu Glu Lys Tyr Glu Glu His Leu

-continued

20	25	30	
Tyr Glu Arg Asp Glu Gly Asp Lys Trp Arg Asn Lys Lys Phe Glu Leu			
35	40	45	
Gly Leu Glu Phe Pro Asn Leu Pro Tyr Tyr Ile Asp Gly Asp Val Lys			
50	55	60	
Leu Thr Gln Ser Met Ala Ile Ile Arg Tyr Ile Ala Asp Lys His Asn			
65	70	75	80
Met Leu Gly Gly Cys Pro Lys Glu Arg Ala Glu Ile Ser Met Leu Glu			
85	90	95	
Gly Ala Val Leu Asp Ile Arg Tyr Gly Val Ser Arg Ile Ala Tyr Ser			
100	105	110	
Lys Asp Phe Glu Thr Leu Lys Val Asp Phe Leu Ser Lys Leu Pro Glu			
115	120	125	
Met Leu Lys Met Phe Glu Asp Arg Leu Cys His Lys Thr Tyr Leu Asn			
130	135	140	
Gly Asp His Val Thr His Pro Asp Phe Met Leu Tyr Asp Ala Leu Asp			
145	150	155	160
Val Val Leu Tyr Met Asp Pro Met Cys Leu Asp Ala Phe Pro Lys Leu			
165	170	175	
Val Cys Phe Lys Lys Arg Ile Glu Ala Ile Pro Gln Ile Asp Lys Tyr			
180	185	190	
Leu Lys Ser Ser Lys Tyr Ile Ala Trp Pro Leu Gln Gly Trp Gln Ala			
195	200	205	
Thr Phe Gly Gly Asp His Pro Pro Lys Ser Asp Leu Val Pro Arg			
210	215	220	
Gly Ser Pro Gly Ile Ser Gly Gly Gly Gly Ile Leu Asp Ser Met			
225	230	235	240
Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys Ser Thr Ala			
245	250	255	
Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val			
260	265	270	
Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr			
275	280	285	
Asn Leu			
290			

<210> SEQ ID NO 29
<211> LENGTH: 972
<212> TYPE: DNA
<213> ORGANISM: *Salmonella dublin*

<400> SEQUENCE: 29

atgcgggggtt ctccatcatca tcatcatcat ggttatggcta gcatgactgg tggacagcaa 60
atgggtcggg atctgtacga ccatgacgat aaggatccga tggcacaagt cattaataca 120
aacagcctgt cgctgttgac ccagaataac ctgaacaaat ctcagtcctc acttagttcc 180
gctattgagc gtctgtccctc tggctgcgt atcaacacgacg cggaaagacgta tgccggcaggc 240
caggcgattt ctaaccgctt cacttctaat atcaaaggcc tgactcaggc ttcccgtaac 300
gctaacgacg gcatttctat tgccgcagacc actgaagggtg cgctgaatgaa atcaacaac 360
aacctgcacg gtgtgcgtga gttgtctgtt caggccacta acgggactaa ctctgattcc 420
gtatctgaaat ctatccagga taaaattcag caacgtctgg aagaaatcga tgcgcgtttct 480
aatcagactc aatttaacgg tggttaaagtc ctctctcagg acaaccagat qaaaatccag 540

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gttggtgcta acgatggtga aaccattacc atcgatctgc	aaaaaattga tgtgaaaagc	600
cttggcccta tccccggaat ttccgggtggt ggtgggtggaa	ttcttagactc catgggtaca	660
ttaatcaatg aagacgctgc cgcagccaag aaaagtacct	ctaaccact ggcttcaatt	720
gattctgcatttgtcaaagt ggacgcagg ttcttctc tgggggcaat	tcaaaaccgt	780
tttgattcag ccattaccaa ccttggcaat acggtaacca	atctgaactc cgcgcttagc	840
cgtatcagaatgctgacta tgcaacggaa gtttctaata	tgtctaaagc gcagattctg	900
cagcaggctgtacttccgt tctggcgcag gctaaccagg	ttccgcaaaa cgtcctctct	960
ttactgcgtt ag		972

<210> SEQ ID NO 30

<211> LENGTH: 323

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 30

Met Arg Gly Ser His His His His His His Gly	Met Ala Ser Met Thr	
1 5 10 15		
Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Asp Lys Asp		
20 25 30		
Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln		
35 40 45		
Asn Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg		
50 55 60		
Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly		
65 70 75 80		
Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln		
85 90 95		
Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu		
100 105 110		
Gly Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu		
115 120 125		
Ser Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser		
130 135 140		
Ile Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser		
145 150 155 160		
Asn Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln		
165 170 175		
Met Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp		
180 185 190		
Leu Gln Lys Ile Asp Val Lys Ser Leu Gly Leu Ile Pro Gly Ile Ser		
195 200 205		
Gly Gly Gly Gly Ile Leu Asp Ser Met Gly Thr Leu Ile Asn Glu		
210 215 220		
Asp Ala Ala Ala Ala Lys Lys Ser Thr Ala Asn Pro Leu Ala Ser Ile		
225 230 235 240		
Asp Ser Ala Leu Ser Lys Val Asp Ala Val Arg Ser Ser Leu Gly Ala		
245 250 255		
Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn Leu Gly Asn Thr Val		
260 265 270		
Thr Asn Leu Asn Ser Ala Arg Ser Arg Ile Glu Asp Ala Asp Tyr Ala		
275 280 285		
Thr Glu Val Ser Asn Met Ser Lys Ala Gln Ile Leu Gln Gln Ala Gly		

290	295	300
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Thr Ser Val Leu Ala Gln Ala Asn Gln Val Pro Gln Asn Val Leu Ser
305 310 315 320

Leu Leu Arg

<210> SEQ ID NO 31

<211> LENGTH: 813

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 31

atgcgggggtt ctcatcatca tcacatcatcat ggtatggcta gcatgactgg tggacagcaa	60
atgggtcggg atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggc	120
ctgactcagg cttcccgtaa cgctaaccgac ggcatattcta ttgcgcagac cactgaaggt	180
gctgctgaatg aaatcaacaa caaacctgcag cgtgtgcgtg agttgtctgt tcaggccact	240
aacggggacta actctgattc cgatctgaaa tctatccagg atgaaattca gcaacgtctg	300
gaagaaaatcg atcgcgttcc taatcagact caatttaacg gtgttaaagt cctctctcag	360
gacaaccaga tgaaaatcca gggtgggtgt aacgatgggt aaaccattac catcgatctg	420
caaaaaattg atgtgaaaag ccttggcctt atccccggaa tttccgggtgg tgggtgggaa	480
attcttagact ccatgggtac attaatcaat gaagacgctg ccgcagccaa gaaaagtacc	540
gtctaaccac tggcttcaat tgattctgca ttgtcaaaag tggacgcagt tcgttcttct	600
ctggggggcaa ttcaaaaccg ttttgattca gccattacca accttggcaa tacggtaacc	660
aatctgaact ccgcgcgtag ccgtatcgaa gatgctgact atgcaacgga agtttctaat	720
atgtctaaag cgcaaggattct gcagcaggct ggtacttccg ttctggcgcgaa ggctaaccag	780
gttccgcaaa acgttctctc tttactgcgt tag	813

<210> SEQ ID NO 32

<211> LENGTH: 270

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 32

Met Arg Gly Ser His His His His His Gly Met Ala Ser Met Thr	
1 5 10 15	

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp	
20 25 30	

Pro Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala Ser Arg Asn Ala	
35 40 45	

Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly Ala Leu Asn Glu	
50 55 60	

Ile Asn Asn Asn Leu Gln Arg Val Arg Glu Leu Ser Val Gln Ala Thr	
65 70 75 80	

Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser Ile Gln Asp Glu Ile	
85 90 95	

Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser Asn Gln Thr Gln Phe	
100 105 110	

Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln Met Lys Ile Gln Val	
115 120 125	

Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp Leu Gln Lys Ile Asp	
130 135 140	

Val Lys Ser Leu Gly Leu Ile Pro Gly Ile Ser Gly Gly Gly Gly

-continued

145	150	155	160
Ile Leu Asp Ser Met Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Ala			
165	170	175	
Lys Lys Ser Thr Ala Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser			
180	185	190	
Lys Val Asp Ala Val Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe			
195	200	205	
Asp Ser Ala Ile Thr Asn Leu Gly Asn Thr Val Thr Asn Leu Asn Ser			
210	215	220	
Ala Arg Ser Arg Ile Glu Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn			
225	230	235	240
Met Ser Lys Ala Gln Ile Leu Gln Gln Ala Gly Thr Ser Val Leu Ala			
245	250	255	
Gln Ala Asn Gln Val Pro Gln Asn Val Leu Ser Leu Leu Arg			
260	265	270	

<210> SEQ ID NO 33

<211> LENGTH: 951

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 33

atgcgggggtt ctcatcatca tcatacatcat ggtatggcta gcatgactgg tggacagcaa 60
 atgggtcggg atctgtacga cgatgacgat aaggatccga tggcacaagt cattaataca 120
 aacagectgt cgctgttgac ccagaataac ctgaacaaat ctcagtcctc actgagttcc 180
 gctattgagc gtctgtcctc tggctctgcgt atcaacagcg cgaaagacga tgccggcaggc 240
 caggcgattt ctaaccgcctt cacttctaat atcaaaggcc tgactcaggc ttcccgtaac 300
 gctaacgacg gcatttctat tgccgacacc actgaagggtg cgctgaatga aatcaacaac 360
 aacctgcagc gtgtgcgtga gttgtctgtt cagggcacta acgggactaa ctctgatcc 420
 gatctgaaat ctatccagga tgaatttcag caacgtctgg aagaaaatcga tcgcgttct 480
 aatcagactc aatttaacgg tggtaaagtc ctctctcagg acaaccagat gaaaatccag 540
 gttggtgcta acgatggtga aaccattacc atcgatctgc aaaaaatttat cccggaaatt 600
 tccgggtgggt gtgggtgaat tcttagactcc atgggtacat taatcaatga agacgctgcc 660
 gcagccaaga aaagtaccgc taacccactg gcttcaattt attctgcatt gtcaaaagtgc 720
 gagcgcgttc gttttctctt gggggcaatt caaaaccgtt ttgattcagc cattaccaac 780
 ctggcaata cggtaaccaa tctgaactcc gcgcgtagcc gtatcgaaga tgctgactat 840
 gcaacggaaat tttctaatat gtctaaagcg cagattctgc agcaggctgg tactccgtt 900
 ctggcgagg ctaaccaggt tccgcaaaac gtcctctctt tactgcgtta g 951

<210> SEQ ID NO 34

<211> LENGTH: 316

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 34

Met Arg Gly Ser His His His His His His Gly	Met Ala Ser Met Thr		
1	5	10	15
Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Asp Lys Asp			
20	25	30	
Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln			
35	40	45	

-continued

Asn Asn Leu Asn Lys Ser Gln Ser Ser Ser Leu Ser Ser Ala Ile Glu Arg
 50 55 60
 Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
 65 70 75 80
 Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
 85 90 95
 Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu
 100 105 110
 Gly Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu
 115 120 125
 Ser Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser
 130 135 140
 Ile Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser
 145 150 155 160
 Asn Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln
 165 170 175
 Met Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp
 180 185 190
 Leu Gln Lys Ile Ile Pro Gly Ile Ser Gly Gly Gly Gly Ile Leu
 195 200 205
 Asp Ser Met Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys
 210 215 220
 Ser Thr Ala Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val
 225 230 235 240
 Asp Ala Val Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser
 245 250 255
 Ala Ile Thr Asn Leu Gly Asn Thr Val Thr Asn Leu Asn Ser Ala Arg
 260 265 270
 Ser Arg Ile Glu Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser
 275 280 285
 Lys Ala Gln Ile Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala
 290 295 300
 Asn Gln Val Pro Gln Asn Val Leu Ser Leu Leu Arg
 305 310 315

<210> SEQ ID NO 35

<211> LENGTH: 792

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 35

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atgcgggggtt ctcatcatca tcatcatcat ggtatggcta gcatgactgg tggacagcaa 60
atgggtcggtt atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggc 120
ctgactcagg cttcccgtaa cgctaaccgac ggcatttcta ttgcgcagac cactgaaggt 180
ggcgtgaatg aaatcaacaa caacctgcag cgtgtgcgtg agttgtctgt tcaggccact 240
aacgggacta actctgattc cgatctgaaa tctatccagg atgaaattca gcaacgtctg 300
gaagaaaatcg atcgcgtttc taatcagact caatttaacg gtgttaaagt cctctctcag 360
gacaaccaga tgaaaatcca gggtggtgct aacgatggtg aaaccattac catcgatctg 420
caaaaaatta tccccggaat ttccgggtgtt ggtgggtggaa ttcttagactc catgggtaca 480
ttaatcaatg aagacgctgc cgccagccaag aaaagtaccg ctaaccact ggcttcaatt 540
gattctgcatttgtcaaaatggacgcagtt cggttcttc tgggggcaat tcaaaaccgt 600

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tttgcgtttt ttatccatcc tttttttttt tttttttttt	660
cgtatcgaaat gttttttttt ttatccatcc tttttttttt	720
cggccggccg tttttttttt ttatccatcc tttttttttt	780
ttacttgcgtt ag	792

<210> SEQ ID NO 36
<211> LENGTH: 263
<212> TYPE: PRT
<213> ORGANISM: *Salmonella dublin*

<400> SEQUENCE: 36

Met	Arg	Gly	Ser	His	His	His	His	His	Gly	Met	Ala	Ser	Met	Thr
1				5					10				15	

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Asp Lys Asp
20 25 30

Pro Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala Ser Arg Asn Ala
35 40 45

Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly Ala Leu Asn Glu
50 55 60

Ile	Asn	Asn	Asn	Leu	Gln	Arg	Val	Arg	Glu	Leu	Ser	Val	Gln	Ala	Thr
65					70					75					80

Asn	Gly	Thr	Asn	Ser	Asp	Ser	Asp	Leu	Lys	Ser	Ile	Gln	Asp	Glu	Ile	
														85	90	95

Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser Asn Gln Thr Gln Phe
 100 105 110

Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln Met Lys Ile Gln Val
115 120 125

Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp Leu Gln Lys Ile Ile
 130 135 140

Pro Gly Ile Ser Gly Gly Gly Gly Gly Ile Leu Asp Ser Met Gly Thr
145 150 155 160

Leu Ile Asn Glu Asp Ala Ala Ala Ala Lys Lys Ser Thr Ala Asn Pro
165 170 175

Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val Arg Ser
180 185 190

Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn Leu
195 200 205

Gly Asn Thr Val Thr Asn Leu Asn Ser Ala Arg Ser Arg Ile Glu Asp
S10 S15 S20

Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Lys Ala Gln Ile Leu

Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val Pro Gln

Asn Val Leu Ser Leu Leu Arg

<211> LENGTH: 807
<212> TYPE: DNA
<213> ORGANISM: *Salmonella dublin*

<400> SEQUENCE: 37

```
atgcggggtt ctccatcatca tcatcatcat ggttatggcta gcatgactgg tggacagcaa 60  
atgggtcggg atctgtacga cggatgacgat aaggatccga tggcacaagt cattaataca 120
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aacagcctgt cgctggtgac ccagaataac ctgaacaaat ctcagtcctc actgagttcc      180
getattgagc gtctgtcctc tggctgcgt atcaacagcg cgaaagacga tgccggcaggc      240
caggcgattt gtaaccgctt cacttctaat atcaaaggcc tgactcaggc ttcccgtaac      300
gctaaccgacg gcatttctat tgccgagacc actgaaggtg cgctgaatga aatcaacaac      360
aacctgcagc gtgtgcgtga gttgtctgtt cagggccacta acgggactaa ctctgattcc      420
gatctgaaat ctatccagga tggaaattcg caacgtctgg aagaaatcga tcgcgttct      480
aatcagactc aatttaacgg tggtaaagtct ctctctcagg acaaccagat gaaaatccag      540
gttgggtctt aacgatggtga aaccattacc atcgatctgc aaaaaatttg tggtaaaagc      600
cttggcctta tccccggaaat ttccgggttgtt ggtgggtggaa ttctagactc catgggtaca      660
ttaatcaatg aagacgctgc cgcagccaag aaaagtaccg ctaaccact ggcttcaatt      720
gattctgcatt tggtaaaagt ggacgcagtt cgttcttctc tgggggcaat tcaaaaccgt      780
tttgatttag ccattaccaa ccttttag                                         807

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<210> SEQ ID NO 38
<211> LENGTH: 268
<212> TYPE: PRT
<213> ORGANISM: *Salmonella dublin*

<400> SEQUENCE: 38

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Met Arg Gly Ser His His His His His His Gly Met Ala Ser Met Thr
1           5           10          15

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp
20          25           30

Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln
35           40           45

Asn Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg
50           55           60

Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
65           70           75           80

Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
85           90           95

Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu
100          105          110

Gly Ala Leu Asn Glu Ile Asn Asn Asn Leu Gln Arg Val Arg Glu Leu
115          120          125

Ser Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser
130          135          140

Ile Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser
145          150          155          160

Asn Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln
165          170          175

Met Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp
180          185          190

Leu Gln Lys Ile Asp Val Lys Ser Leu Gly Leu Ile Pro Gly Ile Ser
195          200          205

Gly Gly Gly Gly Ile Leu Asp Ser Met Gly Thr Leu Ile Asn Glu
210          215          220

Asp Ala Ala Ala Lys Lys Ser Thr Ala Asn Pro Leu Ala Ser Ile
225          230          235          240

Asp Ser Ala Leu Ser Lys Val Asp Ala Val Arg Ser Ser Leu Gly Ala

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245	250	255
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Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn Leu
260 265

<210> SEQ ID NO 39

<211> LENGTH: 786

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 39

atgcggggtt	ctcatcatca	tcatcatcat	ggtatggcta	gcatgactgg	tggacagcaa	60
atgggtcggg	atctgtacga	cgatgacgat	aaggatccga	tggcacaagt	cattaataca	120
aacagectgt	cgctgttgac	ccagaataac	ctgaacaaat	ctcagtcctc	actgagttcc	180
getattgagc	gtctgtcttc	tggctctgcgt	atcaacagcg	cgaaaagacga	tgcggcaggc	240
caggcgattg	ctaaccgctt	cacttctaat	atcaaaggcc	tgactcaggc	ttcccgtaac	300
gctaacgacg	gcatttctat	tgccgagacc	actgaaggtg	cgctgaatga	aatcaacaac	360
aacctgcagc	gtgtgcgtga	gttgcgtgtt	caggccacta	acgggactaa	ctctgattcc	420
gatctgaaat	otatccagga	tgaaattcag	caacgtctgg	aagaaatcga	tgcgtttct	480
aatcagactc	aatttaacgg	tgttaaagtc	ctctctcagg	acaaccagat	aaaaatccag	540
gttgggtgcta	acgatggtga	aaccattacc	atcgatctgc	aaaaaattat	cccggaatt	600
tccgggtggtg	gtgggtggat	tctagactcc	atgggtacat	taatcaatga	agacgctgcc	660
gcagccaaga	aaagtaccgc	taaccactg	gcttcaattg	attctgcatt	gtcaaaagtg	720
gacgcagttc	gttcttctct	gggggcaatt	caaaaccgtt	ttgattcagc	cattaccaac	780
cttttag						786

<210> SEQ ID NO 40

<211> LENGTH: 261

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 40

Met	Arg	Gly	Ser	His	His	His	His	His	His	Gly	Met	Ala	Ser	Met	Thr
1											10				15

Gly	Gly	Gln	Gln	Met	Gly	Arg	Asp	Leu	Tyr	Asp	Asp	Asp	Asp	Lys	Asp
				20				25						30	

Pro	Met	Ala	Gln	Val	Ile	Asn	Thr	Asn	Ser	Leu	Ser	Leu	Leu	Thr	Gln
				35				40						45	

Asn	Asn	Leu	Asn	Lys	Ser	Gln	Ser	Ser	Leu	Ser	Ser	Ala	Ile	Glu	Arg
				50				55						60	

Leu	Ser	Ser	Gly	Leu	Arg	Ile	Asn	Ser	Ala	Lys	Asp	Asp	Ala	Ala	Gly
				65				70						80	

Gln	Ala	Ile	Ala	Asn	Arg	Phe	Thr	Ser	Asn	Ile	Lys	Gly	Leu	Thr	Gln
						85				90				95	

Ala	Ser	Arg	Asn	Ala	Asn	Asp	Gly	Ile	Ser	Ile	Ala	Gln	Thr	Thr	Glu
				100				105						110	

Gly	Ala	Leu	Asn	Glu	Ile	Asn	Asn	Leu	Gln	Arg	Val	Arg	Glu	Leu	
				115				120						125	

Ser	Val	Gln	Ala	Thr	Asn	Gly	Thr	Asn	Ser	Asp	Ser	Asp	Leu	Lys	Ser
				130				135						140	

Ile	Gln	Asp	Glu	Ile	Gln	Gln	Arg	Leu	Glu	Glu	Ile	Asp	Arg	Val	Ser
				145				150						155	

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Asn	Gln	Thr	Gln	Phe	Asn	Gly	Val	Lys	Val	Leu	Ser	Gln	Asp	Asn	Gln
165							170							175	

Met	Lys	Ile	Gln	Val	Gly	Ala	Asn	Asp	Gly	Glu	Thr	Ile	Thr	Ile	Asp
180							185							190	

Leu	Gln	Lys	Ile	Ile	Pro	Gly	Ile	Ser	Gly	Gly	Gly	Gly	Ile	Leu	
195							200					205			

Asp	Ser	Met	Gly	Thr	Leu	Ile	Asn	Glu	Asp	Ala	Ala	Ala	Lys	Lys	
210							215					220			

Ser	Thr	Ala	Asn	Pro	Leu	Ala	Ser	Ile	Asp	Ser	Ala	Leu	Ser	Lys	Val
225							230					235			240

Asp	Ala	Val	Arg	Ser	Ser	Leu	Gly	Ala	Ile	Gln	Asn	Arg	Phe	Asp	Ser
245							250					255			

Ala	Ile	Thr	Asn	Leu											
							260								

<210> SEQ ID NO 41

<211> LENGTH: 849

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 41

atgcgggggtt	ctcatcatca	tcatcatcat	ggtatggcta	gcatgactgg	tggacagcaa		60
atgggtcggg	atctgtacga	cgatgacgat	aaggatccga	tggcacaagt	cattaataca		120
aacagectgt	cgctgttgac	ccagaataac	ctgaacaaat	ctcagtcctc	actgagttcc		180
gttattgagc	gtctgtccctc	tggctctgctgt	atcaacagcg	cgaaagacga	tgccggcaggc		240
caggcgattg	ctaaccgctt	cacttctaat	atcaaaggcc	tgactcaggc	ttcccgtaac		300
gctaaccgacg	gcatttctat	tgccgagacc	actgaagggt	cgctgaatga	aatcaacaaac		360
aacctgcagc	gtgtgcgtga	gttgcgttgc	caggccacta	acgggactaa	ctctgattcc		420
gatctgaaat	ctatccagga	tgaaattcag	caacgtctgg	aagaaaatcga	tcgcgtttct		480
aatcagatcc	cgggaaatttc	cgggtgggt	ggtggaaattc	tagactccat	gggtacattta		540
atcaatgaag	acgctgccgc	agccaagaaa	agtaccgcta	acccactggc	ttcaatttgc		600
tctgcattgt	caaaaatggc	cgcagtttgt	tcttctctgg	gggcaattca	aaaccgtttt		660
gattcagcca	ttaccaaccc	tggcaatacg	gtaaccaatc	tgaactccgc	gcgttagccgt		720
atcgaagatg	ctgactatgc	aacgaaagtt	tctaataatgt	ctaaagcgc	gattctgcag		780
caggctggta	cttccgttct	ggcgcaggct	aaccaggttc	cgcaaaacgt	cctctttta		840
ctgcgttag							849

<210> SEQ ID NO 42

<211> LENGTH: 282

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 42

Met	Arg	Gly	Ser	His	His	His	His	His	Gly	Met	Ala	Ser	Met	Thr	
1							5							15	

Gly	Gly	Gln	Gln	Met	Gly	Arg	Asp	Leu	Tyr	Asp	Asp	Asp	Lys	Asp	
20							25						30		

Pro	Met	Ala	Gln	Val	Ile	Asn	Thr	Asn	Ser	Leu	Ser	Leu	Leu	Thr	Gln
35							40					45			

Asn	Asn	Leu	Asn	Lys	Ser	Gln	Ser	Ser	Leu	Ser	Ser	Ala	Ile	Glu	Arg
50							55					60			

-continued

Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
 65 70 75 80
 Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
 85 90 95
 Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu
 100 105 110
 Gly Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu
 115 120 125
 Ser Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser
 130 135 140
 Ile Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser
 145 150 155 160
 Asn Gln Ile Pro Gly Ile Ser Gly Gly Gly Gly Ile Leu Asp Ser
 165 170 175
 Met Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys Ser Thr
 180 185 190
 Ala Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala
 195 200 205
 Val Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile
 210 215 220
 Thr Asn Leu Gly Asn Thr Val Thr Asn Leu Asn Ser Ala Arg Ser Arg
 225 230 235 240
 Ile Glu Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Lys Ala
 245 250 255
 Gln Ile Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln
 260 265 270
 Val Pro Gln Asn Val Leu Ser Leu Leu Arg
 275 280

<210> SEQ ID NO 43
 <211> LENGTH: 690
 <212> TYPE: DNA
 <213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 43

```

atgcggggttt ctcatcatca tcatacatcat ggtatggcta gcatgactgg tggacagcaa      60
atggggcgggg atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggc      120
ctgactcagg cttcccgtaa cgctaaccgac ggcatttcta ttgcgcagac cactgaaggt      180
gcgcgtgaatg aaatcaacaa caaacctgcag cgtgtgcgtg agttgtctgt tcaggccact      240
aacggggacta actctgattc cgatctgaaa tctatccagg atgaaattca gcaacgtctg      300
gaagaaaatcg atcgcgtttc taatcagatc ccgggaattt ccgggtgggg tggtgaaatt      360
ctagactcca tgggtacatt aatcaatgaa gacgcgtcccg cagccaagaa aagtaccgct      420
aaccccactgg ottcaattga ttctgcatttgc tcaaaagtgg acgcagttcg ttcttctctg      480
ggggcaatttca aaaaccgttt tgattcagcc attaccaacc ttggcaatac ggtaaccaat      540
ctgaactccg cgccgtagccg tattcaaatgat gctgactatg caacggaaat ttcttaatatg      600
tctaaagcgc agattctgca gcaggctgtt acttccgttc tggcgcagac taaccaggtt      660
ccgcacaaacg tcctctctttt actgcgttag                                         690
  
```

<210> SEQ ID NO 44
 <211> LENGTH: 229
 <212> TYPE: PRT
 <213> ORGANISM: Salmonella dublin

-continued

<400> SEQUENCE: 44

Met	Arg	Gly	Ser	His	His	His	His	His	Gly	Met	Ala	Ser	Met	Thr
1				5			10			15				

Gly	Gly	Gln	Gln	Met	Gly	Arg	Asp	Leu	Tyr	Asp	Asp	Asp	Lys	Asp
				20		25			30					

Pro	Phe	Thr	Ser	Asn	Ile	Lys	Gly	Leu	Thr	Gln	Ala	Ser	Arg	Asn	Ala
				35		40			45						

Asn	Asp	Gly	Ile	Ser	Ile	Ala	Gln	Thr	Glu	Gly	Ala	Leu	Asn	Glu
	50				55			60						

Ile	Asn	Asn	Leu	Gln	Arg	Val	Arg	Glu	Leu	Ser	Val	Gln	Ala	Thr
65				70			75		80					

Asn	Gly	Thr	Asn	Ser	Asp	Ser	Asp	Leu	Lys	Ser	Ile	Gln	Asp	Glu	Ile
	85				90			95							

Gln	Gln	Arg	Leu	Glu	Glu	Ile	Asp	Arg	Val	Ser	Asn	Gln	Ile	Pro	Gly
	100				105			110							

Ile	Ser	Gly	Gly	Gly	Gly	Ile	Leu	Asp	Ser	Met	Gly	Thr	Leu	Ile
	115				120			125						

Asn	Glu	Asp	Ala	Ala	Ala	Ala	Lys	Lys	Ser	Thr	Ala	Asn	Pro	Leu	Ala
	130				135			140							

Ser	Ile	Asp	Ser	Ala	Leu	Ser	Lys	Val	Asp	Ala	Val	Arg	Ser	Ser	Leu
145					150			155			160				

Gly	Ala	Ile	Gln	Asn	Arg	Phe	Asp	Ser	Ala	Ile	Thr	Asn	Leu	Gly	Asn
	165				170			175							

Thr	Val	Thr	Asn	Leu	Asn	Ser	Ala	Arg	Ser	Arg	Ile	Glu	Asp	Ala	Asp
	180				185			190							

Tyr	Ala	Thr	Glu	Val	Ser	Asn	Met	Ser	Lys	Ala	Gln	Ile	Leu	Gln	Gln
	195				200			205							

Ala	Gly	Thr	Ser	Val	Leu	Ala	Gln	Ala	Asn	Gln	Val	Pro	Gln	Asn	Val
	210				215			220							

Leu	Ser	Leu	Leu	Arg											
	225														

<210> SEQ ID NO 45

<211> LENGTH: 684

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 45

atgcgggggtt	ctcatcatca	tcatcatcat	ggtatggcta	gcatgactgg	tggacagcaa	60
-------------	------------	------------	------------	------------	------------	----

atgggtcgaaa	atctgtacga	cgtatgacgt	aaggatccga	tggcacaagt	cattaataca	120
-------------	------------	------------	------------	------------	------------	-----

aacagecctgt	cgcgttgac	ccagaataac	ctgaacaaat	ctcagtcctc	actgagttcc	180
-------------	-----------	------------	------------	------------	------------	-----

gtatttgagc	gtctgtcctc	tggctctgcgt	atcaacagcg	cgaaagacga	tgcggcaggg	240
------------	------------	-------------	------------	------------	------------	-----

caggcgattg	ctaaccgctt	cacttctaat	atcaaaggcc	tgactcaggc	ttccccgttaac	300
------------	------------	------------	------------	------------	--------------	-----

gtctgtcctat	tgcgacgacc	actgaagggt	cgctgaatga	aatcaacaaac		360
-------------	------------	------------	------------	-------------	--	-----

accctgcagc	gtgtgcgtga	gttgtctgtt	caggccacta	acgggactaa	ctctgatcc	420
------------	------------	------------	------------	------------	-----------	-----

gatctgaaat	ctatccagga	tgaaattcag	caacgtctgg	aagaaatcga	tgcgtttct	480
------------	------------	------------	------------	------------	-----------	-----

aatcagatcc	cgggaatttc	cggtggtgg	ggtggaaattc	tagactccat	gggtacatta	540
------------	------------	-----------	-------------	------------	------------	-----

atcaatgaag	acgctgccgc	agccaagaaa	agtaccgcta	acccactggc	ttcaatttgc	600
------------	------------	------------	------------	------------	------------	-----

tctgcattgt	caaaaagtgg	cgcagttcgt	tcttctctgg	gggcaattca	aaaccgtttt	660
------------	------------	------------	------------	------------	------------	-----

gattcagcca	ttaccaacct	tttag				684
------------	------------	-------	--	--	--	-----

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<210> SEQ_ID NO 46
<211> LENGTH: 227
<212> TYPE: PRT
<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 46

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Met Arg Gly Ser His His His His His Gly Met Ala Ser Met Thr
      5           10          15

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp
     20            25          30

Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln
      35            40          45

Asn Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg
      50            55          60

Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
      65            70          75          80

Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
      85            90          95

Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu
     100           105         110

Gly Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu
     115           120         125

Ser Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser
     130           135         140

Ile Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser
     145           150         155         160

Asn Gln Ile Pro Gly Ile Ser Gly Gly Gly Gly Ile Leu Asp Ser
     165           170         175

Met Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Lys Lys Ser Thr
     180           185         190

Ala Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala
     195           200         205

Val Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile
     210           215         220

Thr Asn Leu
     225

```

<210> SEQ_ID NO 47
<211> LENGTH: 525
<212> TYPE: DNA
<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 47

```

atgcggggtt ctcatcatca tcatacatcat ggtatggcta gcatgactgg tggacagcaa    60
atgggtcggtt atctgtacga cgatgacgat aaggatccgt tcacttctaa tatcaaaggc    120
ctgactcagg cttcccgtaa cgctaaccgac ggcatttcta ttgcgcagac cactgaaggt    180
gcgctgaatg aaatcaacaa caacctgcag cgtgtgcgtg agttgtctgt tcaggccact    240
aacgggacta actctgattc cgatctgaaa tctatccagg atgaaattca gcaacgtctg    300
gaagaaaatcg atcgcgtttc taatcagatc ccggaaattt ccgggtgggg tggtggatt    360
ctagactcca tgggtacatt aatcaatgaa gacgctgccg cagccaagaa aagtaccgct    420
aacccactgg cttcaattga ttctgcattt tcaaaaagtgg acgcagttcg ttcttcttg    480
ggggcaattc aaaaccgttt tgattcagcc attaccaacc ttttag    525

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<210> SEQ_ID NO 48
<211> LENGTH: 174
<212> TYPE: PRT
<213> ORGANISM: *Salmonella dublin*

<400> SEQUENCE: 48

Met	Arg	Gly	Ser	His	His	His	His	His	Gly	Met	Ala	Ser	Met	Thr
1				5			10			15				

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp
20 25 30

Pro	Phe	Thr	Ser	Asn	Ile	Lys	Gly	Leu	Thr	Gln	Ala	Ser	Arg	Asn	Ala
35					40				45						

Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly Ala Leu Asn Glu
50 55 60

Ile	Asn	Asn	Asn	Leu	Gln	Arg	Val	Arg	Glu	Leu	Ser	Val	Gln	Ala	Thr
65				70			75		80						

Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser Ile Gln Asp Glu Ile
85 90 95

Gln	Gln	Arg	Leu	Glu	Glu	Ile	Asp	Arg	Val	Ser	Asn	Gln	Ile	Pro	Gly
100				105				110							

Ile Ser Gly Gly Gly Ile Leu Asp Ser Met Gly Thr Leu Ile
115 120 125

Asn	Glu	Asp	Ala	Ala	Ala	Lys	Lys	Ser	Thr	Ala	Asn	Pro	Leu	Ala	
130					135			140							

Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val Arg Ser Ser Leu
145 150 155 160

Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn Leu
165 170

<210> SEQ_ID NO 49
<211> LENGTH: 762
<212> TYPE: DNA
<213> ORGANISM: *Salmonella dublin*

<400> SEQUENCE: 49

atgcgggggtt	ctcatcatca	tcatcatcat	ggtatggcta	gcatgactgg	tggacagcaa	60
atggggtcggg	atctgtacga	cgtatcgat	aaggatccga	tggcacaagt	cattaataca	120
aacagcctgt	cgctgttgac	ccagaataac	ctgaacaaat	ctcagtcctc	actgagttcc	180
gttattgagc	gtctgtcctc	tggctgcgt	atcaacagcg	cgaaagacga	tgcggcaggc	240
caggcgattg	ctaaccgctt	cacttctaat	atcaaaggcc	tgactcaggc	ttcccgtaac	300
getaacgacg	gcatttctat	tgcgcagacc	actgaagggt	cgctgaatga	aatcaacaac	360
aacctgcagc	gtgtgcgtga	gttgtctgtt	caggccacta	tcccggaaat	ttccgggtgt	420
ggtgtggaa	ttcttagactc	catgggtaca	ttaatcaatg	aagacgctgc	cgcagccaag	480
aaaagtacct	ctaaccact	ggcttcaatt	gattctgcat	tgtcaaaagt	ggacgcagtt	540
cgttcttc	tggggcaat	tcaaaaccgt	tttgattcag	ccattaccaa	ccttggcaat	600
acggtaacca	atctgaactc	cgcgcgtac	cgtatcgaag	atgctgacta	tgcaacggaa	660
gtttctaata	tgtctaaagc	gcagattctg	cagcaggctg	gtacttccgt	tctggcgcag	720
gctaaccagg	ttccgaaaaa	cgtctctct	ttactgcgtt	ag		762

<210> SEQ_ID NO 50
<211> LENGTH: 253

-continued

<212> TYPE: PRT

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 50

Met Arg Gly Ser His His His His His His Gly Met Ala Ser Met Thr
 1 5 10 15

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp
 20 25 30

Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln
 35 40 45

Asn Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg
 50 55 60

Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
 65 70 75 80

Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
 85 90 95

Ala Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu
 100 105 110

Gly Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu
 115 120 125

Ser Val Gln Ala Thr Ile Pro Gly Ile Ser Gly Gly Gly Gly Ile
 130 135 140

Leu Asp Ser Met Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala Ala Lys
 145 150 155 160

Lys Ser Thr Ala Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys
 165 170 175

Val Asp Ala Val Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp
 180 185 190

Ser Ala Ile Thr Asn Leu Gly Asn Thr Val Thr Asn Leu Asn Ser Ala
 195 200 205

Arg Ser Arg Ile Glu Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met
 210 215 220

Ser Lys Ala Gln Ile Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln
 225 230 235 240

Ala Asn Gln Val Pro Gln Asn Val Leu Ser Leu Leu Arg
 245 250

<210> SEQ ID NO 51

<211> LENGTH: 597

<212> TYPE: DNA

<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 51

atgcgggggtt ctcatcatca tcatacatcat ggtatggcta gcatgactgg tggacagcaa 60
 atgggtcggg atctgtacga cgatgacgat aaggatccga tggcacaagt cattaataca 120
 aacagcctgt cgctgttgac ccagaataac ctgaacaaat ctcagtcctc actgagttcc 180
 gctattgagc gtctgtcctc tggctctgcgt atcaacagcg cgaaagacga tgcggcaggc 240
 caggcgattg otaaccgctt cacttctaat atcaaaggcc tgactcaggc ttcccgtaac 300
 gctaaccgacg gcatttctat tgccgacacc actgaagggt cgctgaatga aatcaacaac 360
 aacctgcagc gtgtgcgtga gttgtctgtt caggccacta tccccggaaat ttccgggttgt 420
 ggtgggtggaa ttcttagactc catgggtaca ttaatcaatg aagacgctgc cgccagccaag 480
 aaaagtaccg otaaccact ggcttcaatt gattctgcat tgtcaaaaagt ggacgcagtt 540

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cgttcttc tggggcaat tcaaaccgt tttgattcag ccattaccaa ccttttag 597

<210> SEQ ID NO 52
<211> LENGTH: 198
<212> TYPE: PRT
<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 52

Met	Arg	Gly	Ser	His	His	His	His	His	Gly	Met	Ala	Ser	Met	Thr
1														
														15

Gly	Gly	Gln	Gln	Met	Gly	Arg	Asp	Leu	Tyr	Asp	Asp	Asp	Lys	Asp
														30
20		25												

Pro	Met	Ala	Gln	Val	Ile	Asn	Thr	Asn	Ser	Leu	Ser	Leu	Leu	Thr	Gln
															45
35		40													

Asn	Asn	Leu	Asn	Lys	Ser	Gln	Ser	Ser	Leu	Ser	Ser	Ala	Ile	Glu	Arg
															60
50		55													

Leu	Ser	Ser	Gly	Leu	Arg	Ile	Asn	Ser	Ala	Lys	Asp	Asp	Ala	Ala	Gly
															80
65		70													

Gln	Ala	Ile	Ala	Asn	Arg	Phe	Thr	Ser	Asn	Ile	Lys	Gly	Leu	Thr	Gln
															95
85		90													

Ala	Ser	Arg	Asn	Ala	Asn	Asp	Gly	Ile	Ser	Ile	Ala	Gln	Thr	Thr	Glu
															110
100		105													

Gly	Ala	Leu	Asn	Glu	Ile	Asn	Asn	Leu	Gln	Arg	Val	Arg	Glu	Leu	
															125
115		120													

Ser	Val	Gln	Ala	Thr	Ile	Pro	Gly	Ile	Ser	Gly	Gly	Gly	Gly	Ile	
															140
130		135													

Leu	Asp	Ser	Met	Gly	Thr	Leu	Ile	Asn	Glu	Asp	Ala	Ala	Ala	Lys	
															160
145		150													

Lys	Ser	Thr	Ala	Asn	Pro	Leu	Ala	Ser	Ile	Asp	Ser	Ala	Leu	Ser	Lys
															175
165		170													

Val	Asp	Ala	Val	Arg	Ser	Ser	Leu	Gly	Ala	Ile	Gln	Asn	Arg	Phe	Asp
															190
180		185													

Ser	Ala	Ile	Thr	Asn	Leu
195					

<210> SEQ ID NO 53
<211> LENGTH: 672
<212> TYPE: DNA
<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 53

atgcggggtt	ctcatcatca	tcatcatcat	ggtatggcta	gcatgactgg	tggacagcaa	60
------------	------------	------------	------------	------------	------------	----

atgggtcggtt	atctgtacga	cgatgacgtt	aaggatccga	tggcacaagt	cattaataca	120
-------------	------------	------------	------------	------------	------------	-----

aacagcctgt	cgctgttgac	ccagaataac	ctgaacaaat	ctcagtcctc	actgagttcc	180
------------	------------	------------	------------	------------	------------	-----

gctattggac	gtctgtccctc	tggctgtcggt	atcaacagcg	cgaaagacga	tgcggcaggc	240
------------	-------------	-------------	------------	------------	------------	-----

caggcgattt	ctaaccgctt	cacttctaat	atcaaaggcc	tgactcagggc	ttcccgtaac	300
------------	------------	------------	------------	-------------	------------	-----

gctaaccgaca	tcccggaaat	ttccgggtgg	ggtgggtggaa	ttcttagactc	catgggtaca	360
-------------	------------	------------	-------------	-------------	------------	-----

ttaatcaatg	aagacgctgc	cgcagccaag	aaaagtaccc	ctaaccact	ggcttcaatt	420
------------	------------	------------	------------	-----------	------------	-----

gattctgcatt	tgtcaaaagt	ggacgcagg	cgttcttc	tggggcaat	tcaaaccgt	480
-------------	------------	-----------	----------	-----------	-----------	-----

tttgattcag	ccattaccaa	ccttggcaat	acggtaaccc	atctgaactc	cgcgcgtac	540
------------	------------	------------	------------	------------	-----------	-----

cgtatcgaag	atgctgacta	tgcaacggaa	gtttctaata	tgtctaaagc	gcagattctg	600
------------	------------	------------	------------	------------	------------	-----

caggcaggctg	gtacttccgt	tctggcgcag	gctaaccagg	ttccgcaaaa	cgtccctct	660
-------------	------------	------------	------------	------------	-----------	-----

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ttactgcgtt ag

672

<210> SEQ ID NO 54
<211> LENGTH: 223
<212> TYPE: PRT
<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 54

Met	Arg	Gly	Ser	His	His	His	His	His	Gly	Met	Ala	Ser	Met	Thr	
1				5			10			15					
Gly	Gly	Gln	Gln	Met	Gly	Arg	Asp	Leu	Tyr	Asp	Asp	Asp	Lys	Asp	
		20				25				30					
Pro	Met	Ala	Gln	Val	Ile	Asn	Thr	Asn	Ser	Leu	Ser	Leu	Leu	Thr	Gln
	35				40				45						
Asn	Asn	Leu	Asn	Lys	Ser	Gln	Ser	Ser	Leu	Ser	Ser	Ala	Ile	Glu	Arg
	50			55			60								
Leu	Ser	Ser	Gly	Leu	Arg	Ile	Asn	Ser	Ala	Lys	Asp	Asp	Ala	Ala	Gly
65				70			75			80					
Gln	Ala	Ile	Ala	Asn	Arg	Phe	Thr	Ser	Asn	Ile	Lys	Gly	Leu	Thr	Gln
	85				90			95							
Ala	Ser	Arg	Asn	Ala	Asn	Asp	Ile	Pro	Gly	Ile	Ser	Gly	Gly	Gly	
	100				105				110						
Gly	Ile	Leu	Asp	Ser	Met	Gly	Thr	Leu	Ile	Asn	Glu	Asp	Ala	Ala	
	115				120			125							
Ala	Lys	Lys	Ser	Thr	Ala	Asn	Pro	Leu	Ala	Ser	Ile	Asp	Ser	Ala	Leu
	130				135			140							
Ser	Lys	Val	Asp	Ala	Val	Arg	Ser	Ser	Leu	Gly	Ala	Ile	Gln	Asn	Arg
145				150			155			160					
Phe	Asp	Ser	Ala	Ile	Thr	Asn	Leu	Gly	Asn	Thr	Val	Thr	Asn	Leu	Asn
	165				170			175							
Ser	Ala	Arg	Ser	Arg	Ile	Glu	Asp	Ala	Asp	Tyr	Ala	Thr	Glu	Val	Ser
	180				185			190							
Asn	Met	Ser	Lys	Ala	Gln	Ile	Leu	Gln	Gln	Ala	Gly	Thr	Ser	Val	Leu
	195				200			205							
Ala	Gln	Ala	Asn	Gln	Val	Pro	Gln	Asn	Val	Leu	Ser	Leu	Leu	Arg	
	210				215			220							

<210> SEQ ID NO 55
<211> LENGTH: 507
<212> TYPE: DNA
<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 55

atgcgggggtt	ctcatacatca	tcatcatcat	ggtatggcta	gcatgactgg	tggacagcaa	60
atgggtcggtt	atctgtacga	cgtatgcgtat	aaggatccga	tggcacaagt	cattaataca	120
aacagccgtt	cgtcttttgcac	ccagaataac	ctgaacaaat	ctcagtcctc	actgagttcc	180
gttatttgatc	gtctgtccctc	tggctgtcggt	atcaacacgcg	cgaaagacga	tgccggcaggc	240
caggcgattt	ctaaccgttt	cacttctaat	atcaaaggcc	tgactcaggc	ttcccgtaac	300
gtctaaccgaca	tcccggttgc	ttcccggttgc	ggtggttgc	ttcttagactc	catgggtaca	360
ttaatcaatg	aagacgctgc	cgcagccaag	aaaagtacct	ctaaccact	ggcttcaatt	420
gattctgcatt	tgtcaaaatgt	ggacgcgttt	cgttcttc	tgggggcatt	tcaaaaccgt	480
tttgattcag	ccatttaccaa	ccttttag				507

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<210> SEQ ID NO 56
<211> LENGTH: 168
<212> TYPE: PRT
<213> ORGANISM: Salmonella dublin

<400> SEQUENCE: 56

```

Met Arg Gly Ser His His His His His Gly Met Ala Ser Met Thr
1           5          10          15

Gly Gly Gln Gln Met Gly Arg Asp Leu Tyr Asp Asp Asp Lys Asp
20          25          30

Pro Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln
35          40          45

Asn Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg
50          55          60

Leu Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly
65          70          75          80

Gln Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln
85          90          95

Ala Ser Arg Asn Ala Asn Asp Ile Pro Gly Ile Ser Gly Gly Gly
100         105         110

Gly Ile Leu Asp Ser Met Gly Thr Leu Ile Asn Glu Asp Ala Ala Ala
115         120         125

Ala Lys Lys Ser Thr Ala Asn Pro Leu Ala Ser Ile Asp Ser Ala Leu
130         135         140

Ser Lys Val Asp Ala Val Arg Ser Ser Leu Gly Ala Ile Gln Asn Arg
145         150         155         160

Phe Asp Ser Ala Ile Thr Asn Leu
165

```

<210> SEQ ID NO 57
<211> LENGTH: 174
<212> TYPE: PRT
<213> ORGANISM: Salmonella enterica

<400> SEQUENCE: 57

```

Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln Asn
1           5          10          15

Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg Leu
20          25          30

Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
35          40          45

Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala
50          55          60

Ser Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly
65          70          75          80

Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu Ser
85          90          95

Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Lys Ser Ile
100         105         110

Gln Asp Glu Ile Gln Gln Arg Leu Glu Glu Ile Asp Arg Val Ser Asn
115         120         125

Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gln Asp Asn Gln Met
130         135         140

Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asp Leu
145         150         155         160

Gln Lys Ile Asp Val Lys Ser Leu Gly Leu Asp Gly Phe Asn

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165

170

<210> SEQ ID NO 58
<211> LENGTH: 189
<212> TYPE: PRT
<213> ORGANISM: *Pseudomonas aeruginosa*

<400> SEQUENCE: 58

```

Met Ala Leu Thr Val Asn Thr Asn Ile Ala Ser Leu Asn Thr Gln Arg
1           5          10          15

Asn Leu Asn Ala Ser Ser Asn Asp Leu Asn Thr Ser Leu Gln Arg Leu
20          25          30

Thr Thr Gly Tyr Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Leu
35          40          45

Gln Ile Ser Asn Arg Leu Ser Asn Gln Ile Ser Gly Leu Asn Val Ala
50          55          60

Thr Arg Asn Ala Asn Asp Gly Ile Ser Leu Ala Gln Thr Ala Glu Gly
65          70          75          80

Ala Leu Gln Gln Ser Thr Asn Ile Leu Gln Arg Ile Arg Asp Leu Ala
85          90          95

Leu Gln Ser Ala Asn Gly Ser Asn Ser Asp Ala Asp Arg Ala Ala Leu
100         105         110

Gln Lys Glu Val Ala Ala Gln Gln Ala Glu Leu Thr Arg Ile Ser Asp
115         120         125

Thr Thr Thr Phe Gly Gly Arg Lys Leu Leu Asp Gly Ser Phe Gly Thr
130         135         140

Thr Ser Phe Gln Val Gly Ser Asn Ala Tyr Glu Thr Ile Asp Ile Ser
145         150         155         160

Leu Gln Asn Ala Ser Ala Ser Ala Ile Gly Ser Tyr Gln Val Gly Ser
165         170         175

Asn Gly Ala Gly Thr Val Ala Ser Val Ala Gly Thr Ala
180         185

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<210> SEQ ID NO 59
<211> LENGTH: 179
<212> TYPE: PRT
<213> ORGANISM: *Legionella pneumophila*

<400> SEQUENCE: 59

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Met Ala Gln Val Ile Asn Thr Asn Val Ala Ser Leu Thr Ala Gln Arg
1           5          10          15

Asn Leu Gly Val Ser Gly Asn Met Met Gln Thr Ser Ile Gln Arg Leu
20          25          30

Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Leu
35          40          45

Ala Ile Ser Gln Arg Met Thr Ala Gln Ile Arg Gly Met Asn Gln Ala
50          55          60

Val Arg Asn Ala Asn Asp Gly Ile Ser Leu Ala Gln Val Ala Glu Gly
65          70          75          80

Ala Met Gln Glu Thr Thr Asn Ile Leu Gln Arg Met Arg Glu Leu Ser
85          90          95

Val Gln Ala Ala Asn Ser Thr Asn Asn Ser Ser Asp Arg Ala Ser Ile
100         105         110

Gln Ser Glu Ile Ser Gln Leu Lys Ser Glu Leu Glu Arg Ile Ala Gln
115         120         125

Asn Thr Glu Phe Asn Gly Gln Arg Ile Leu Asp Gly Ser Phe Ser Gly

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130 135 140

Ala Ser Phe Gln Val Gly Ala Asn Ser Asn Gln Thr Ile Asn Phe Ser
 145 150 155 160
 Ile Gly Ser Ile Lys Ala Ser Ser Ile Gly Gly Ile Ala Thr Ala Thr
 165 170 175
 Gly Thr Glu

<210> SEQ ID NO 60

<211> LENGTH: 174

<212> TYPE: PRT

<213> ORGANISM: Escherichia coli

<400> SEQUENCE: 60

Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln Asn
 1 5 10 15
 Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg Leu
 20 25 30
 Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
 35 40 45
 Ala Ile Ala Asn Arg Phe Thr Ala Asn Ile Lys Gly Leu Thr Gln Ala
 50 55 60
 Ser Arg Asn Ala Asn Asp Gly Ile Ser Val Ala Gln Thr Thr Glu Gly
 65 70 75 80
 Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu Thr
 85 90 95
 Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Ser Ser Ile
 100 105 110
 Gln Ala Glu Ile Thr Gln Arg Leu Glu Glu Ile Asp Arg Val Ser Glu
 115 120 125
 Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ala Glu Asn Asn Glu Met
 130 135 140
 Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Thr Ile Asn Leu
 145 150 155 160
 Ala Lys Ile Asp Ala Lys Thr Leu Gly Leu Asp Gly Phe Asn
 165 170

<210> SEQ ID NO 61

<211> LENGTH: 173

<212> TYPE: PRT

<213> ORGANISM: Serratia marcescens

<400> SEQUENCE: 61

Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Met Ala Gln Asn
 1 5 10 15
 Asn Leu Asn Lys Ser Gln Ser Ser Leu Gly Thr Ala Ile Glu Arg Leu
 20 25 30
 Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
 35 40 45
 Ala Ile Ser Asn Arg Phe Thr Ala Asn Ile Lys Gly Leu Thr Gln Ala
 50 55 60
 Ser Arg Asn Ala Asn Asp Gly Ile Ser Leu Ala Gln Thr Thr Glu Gly
 65 70 75 80
 Ala Leu Asn Glu Val Asn Asp Asn Leu Gln Asn Ile Arg Arg Leu Thr
 85 90 95
 Val Gln Ala Gln Asn Gly Ser Asn Ser Thr Ser Asp Leu Lys Ser Ile
 100 105 110

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Gln Asp Glu Ile Thr Gln Arg Leu Ser Glu Ile Asn Arg Ile Ser Glu
 115 120 125
 Gln Thr Asp Phe Asn Gly Val Lys Val Leu Ser Ser Asp Gln Lys Leu
 130 135 140
 Thr Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Thr Asp Ile Asp Leu
 145 150 155 160
 Lys Lys Ile Asp Ala Lys Gln Leu Gly Met Asp Thr Phe
 165 170

<210> SEQ ID NO 62
<211> LENGTH: 168
<212> TYPE: PRT
<213> ORGANISM: *Bacillus subtilis*

<400> SEQUENCE: 62

Met Arg Ile Asn His Asn Ile Ala Ala Leu Asn Thr Ser Arg Gln Leu
 1 5 10 15
 Asn Ala Gly Ser Asn Ser Ala Ala Lys Asn Met Glu Lys Leu Ser Ser
 20 25 30
 Gly Leu Arg Ile Asn Arg Ala Gly Asp Asp Ala Ala Gly Leu Ala Ile
 35 40 45
 Ser Glu Lys Met Arg Ser Gln Ile Arg Gly Leu Asp Met Ala Ser Lys
 50 55 60
 Asn Ala Gln Asp Gly Ile Ser Leu Ile Gln Thr Ser Glu Gly Ala Leu
 65 70 75 80
 Asn Glu Thr His Ser Ile Leu Gln Arg Met Ser Glu Leu Ala Thr Gln
 85 90 95
 Ala Ala Asn Asp Thr Asn Thr Asp Ser Asp Arg Ser Glu Leu Gln Lys
 100 105 110
 Glu Met Asp Gln Leu Ala Ser Glu Val Thr Arg Ile Ser Thr Asp Thr
 115 120 125
 Glu Phe Asn Thr Lys Lys Leu Leu Asp Gly Thr Ala Gln Asn Leu Thr
 130 135 140
 Phe Gln Ile Gly Ala Asn Glu Gly Gln Thr Met Ser Leu Ser Ile Asn
 145 150 155 160
 Lys Met Asp Ser Glu Ser Leu Lys
 165

<210> SEQ ID NO 63
<211> LENGTH: 192
<212> TYPE: PRT
<213> ORGANISM: *Listeria monocytogenes*

<400> SEQUENCE: 63

Met Lys Val Asn Thr Asn Ile Ile Ser Leu Lys Thr Gln Glu Tyr Leu
 1 5 10 15
 Arg Lys Asn Asn Glu Gly Met Thr Gln Ala Gln Glu Arg Leu Ala Ser
 20 25 30
 Gly Lys Arg Ile Asn Ser Ser Leu Asp Asp Ala Ala Gly Leu Ala Val
 35 40 45
 Val Thr Arg Met Asn Val Lys Ser Thr Gly Leu Asp Ala Ala Ser Lys
 50 55 60
 Asn Ser Ser Met Gly Ile Asp Leu Leu Gln Thr Ala Asp Ser Ala Leu
 65 70 75 80
 Ser Ser Met Ser Ser Ile Leu Gln Arg Met Arg Gln Leu Ala Val Gln
 85 90 95

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Ser Ser Asn Gly Ser Phe Ser Asp Glu Asp Arg Lys Gln Tyr Thr Ala
 100 105 110

Glu Phe Gly Ser Leu Ile Lys Glu Leu Asp His Val Ala Asp Thr Thr
 115 120 125

Asn Tyr Asn Asn Ile Lys Leu Leu Asp Gln Thr Ala Thr Gly Ala Ala
 130 135 140

Thr Gln Val Ser Ile Gln Ala Ser Asp Lys Ala Asn Asp Leu Ile Asn
 145 150 155 160

Ile Asp Leu Phe Asn Ala Lys Gly Leu Ser Ala Gly Thr Ile Thr Leu
 165 170 175

Gly Ser Gly Ser Thr Val Ala Gly Tyr Ser Ala Leu Ser Val Ala Asp
 180 185 190

<210> SEQ ID NO 64

<211> LENGTH: 174

<212> TYPE: PRT

<213> ORGANISM: Shigella sonnei

<400> SEQUENCE: 64

Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln Asn
 1 5 10 15

Asn Leu Asn Lys Ser Gln Ser Ser Leu Ser Ser Ala Ile Glu Arg Leu
 20 25 30

Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
 35 40 45

Ala Ile Ala Asn Arg Phe Thr Ala Asn Ile Lys Gly Leu Thr Gln Ala
 50 55 60

Ser Arg Asn Ala Asn Asp Gly Ile Ser Val Ala Gln Thr Thr Glu Gly
 65 70 75 80

Ala Leu Ser Glu Ile Asn Asn Leu Gln Arg Ile Arg Glu Leu Ser
 85 90 95

Val Gln Ala Thr Asn Gly Thr Asn Ser Asp Ser Asp Leu Asn Ser Ile
 100 105 110

Gln Asp Glu Ile Thr Gln Arg Leu Ser Glu Ile Asp Arg Val Ser Asn
 115 120 125

Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ala Ser Asp Gln Thr Met
 130 135 140

Lys Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Glu Ile Ala Leu
 145 150 155 160

Asp Lys Ile Asp Ala Lys Thr Leu Gly Leu Asp Asn Phe Ser
 165 170

<210> SEQ ID NO 65

<211> LENGTH: 174

<212> TYPE: PRT

<213> ORGANISM: Edwardsiella tarda

<400> SEQUENCE: 65

Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Met Ala Gln Asn
 1 5 10 15

Asn Leu Asn Lys Ser Gln Ser Ala Leu Gly Thr Ala Ile Glu Arg Leu
 20 25 30

Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
 35 40 45

Ala Ile Ser Asn Arg Phe Thr Ala Asn Ile Asn Gly Leu Thr Gln Ala
 50 55 60

-continued

Ser Arg Asn Ala Asn Asp Gly Ile Ser Leu Ala Gln Thr Thr Glu Gly
 65 70 75 80
 Ala Leu Asn Glu Val Asn Asp Asn Leu Gln Asn Ile Arg Arg Leu Thr
 85 90 95
 Val Gln Ala Gln Asn Gly Ser Asn Ser Ser Asp Leu Gln Ser Ile
 100 105 110
 Gln Asp Glu Ile Thr Gln Arg Leu Ser Glu Ile Asp Arg Ile Ser Gln
 115 120 125
 Gln Thr Asp Phe Asn Gly Val Lys Val Leu Ser Lys Asp Gln Lys Leu
 130 135 140
 Thr Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Asp Ile Asp Leu
 145 150 155 160
 Lys Asn Ile Asn Ala Gln Ser Leu Gly Leu Asp Lys Phe Asn
 165 170

<210> SEQ ID NO 66
<211> LENGTH: 186
<212> TYPE: PRT
<213> ORGANISM: Acidovorax avenae

<400> SEQUENCE: 66

Met Ala Ser Thr Ile Asn Thr Asn Val Ser Ser Leu Thr Ala Gln Arg
 1 5 10 15
 Asn Leu Ser Leu Ser Gln Ser Ser Leu Asn Thr Ser Ile Gln Arg Leu
 20 25 30
 Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Leu
 35 40 45
 Ala Ile Ser Glu Arg Phe Thr Ser Gln Ile Arg Gly Leu Asn Gln Ala
 50 55 60
 Val Arg Asn Ala Asn Asp Gly Ile Ser Leu Ala Gln Thr Ala Glu Gly
 65 70 75 80
 Ala Leu Lys Ser Thr Gly Asp Ile Leu Gln Arg Val Arg Glu Leu Ala
 85 90 95
 Val Gln Ser Ala Asn Ala Thr Asn Ser Ser Gly Asp Arg Lys Ala Ile
 100 105 110
 Gln Ala Glu Val Gly Gln Leu Leu Ser Glu Met Asp Arg Ile Ala Gly
 115 120 125
 Asn Thr Glu Phe Asn Gly Gln Lys Leu Leu Asp Gly Ser Phe Gly Ser
 130 135 140
 Ala Thr Phe Gln Val Gly Ala Asn Ala Asn Gln Thr Ile Thr Ala Thr
 145 150 155 160
 Thr Gly Asn Phe Arg Thr Asn Asn Tyr Gly Ala Gln Leu Thr Ala Ser
 165 170 175
 Ala Ser Gly Ala Ala Thr Ser Gly Ala Ser
 180 185

<210> SEQ ID NO 67
<211> LENGTH: 173
<212> TYPE: PRT
<213> ORGANISM: Yersinia pestis

<400> SEQUENCE: 67

Met Ala Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln Asn Asn
 1 5 10 15
 Leu Asn Lys Ser Gln Ser Ser Leu Gly Thr Ala Ile Glu Arg Leu Ser
 20 25 30

-continued

Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln Ala
 35 40 45

Ile Ala Asn Arg Phe Thr Ser Asn Ile Lys Gly Leu Thr Gln Ala Ala
 50 55 60

Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly Ser
 65 70 75 80

Leu Asn Glu Ile Asn Asn Leu Gln Arg Val Arg Glu Leu Thr Val
 85 90 95

Gln Ala Gln Asn Gly Ser Asn Ser Ser Asp Leu Asp Ser Ile Gln
 100 105 110

Asp Glu Ile Ser Leu Arg Leu Ala Glu Ile Asp Arg Val Ser Asp Gln
 115 120 125

Thr Gln Phe Asn Gly Lys Lys Val Leu Ala Glu Asn Thr Thr Met Ser
 130 135 140

Ile Gln Val Gly Ala Asn Asp Gly Glu Thr Ile Asp Ile Asn Leu Gln
 145 150 155 160

Lys Ile Asp Ser Lys Ser Leu Gly Leu Gly Ser Tyr Ser
 165 170

<210> SEQ ID NO 68

<211> LENGTH: 174

<212> TYPE: PRT

<213> ORGANISM: Photorhabdus luminescens

<400> SEQUENCE: 68

Met Ala Gln Val Ile Asn Thr Asn Ser Leu Ser Leu Leu Thr Gln Asn
 1 5 10 15

Asn Leu Asn Arg Ser Gln Gly Thr Leu Gly Ser Ala Ile Glu Arg Leu
 20 25 30

Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
 35 40 45

Ala Ile Ala Asn Arg Phe Thr Ala Asn Val Arg Gly Leu Thr Gln Ala
 50 55 60

Ala Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly
 65 70 75 80

Ala Leu Asn Glu Ile Asn Thr Asn Leu Gln Arg Ile Arg Glu Leu Thr
 85 90 95

Val Gln Ser Gln Asn Gly Ser Asn Ser Glu Ser Asp Ile Lys Ser Ile
 100 105 110

Gln Glu Glu Val Thr Gln Arg Leu Lys Glu Ile Asp Arg Ile Ser Glu
 115 120 125

Gln Thr Gln Phe Asn Gly Val Arg Val Leu Arg Glu Asp Ser Lys Met
 130 135 140

Thr Ile Gln Val Gly Ala Asn Asp Asn Glu Val Ile Asp Ile Asp Leu
 145 150 155 160

Lys Lys Ile Asp Lys Glu Ala Leu Asn Leu Gly Lys Phe Thr
 165 170

<210> SEQ ID NO 69

<211> LENGTH: 189

<212> TYPE: PRT

<213> ORGANISM: Rhodobacter sphaeroides

<400> SEQUENCE: 69

Met Thr Thr Ile Asn Thr Asn Ile Gly Ala Ile Ala Ala Gln Ala Asn
 1 5 10 15

-continued

Met Thr Lys Val Asn Asp Gln Phe Asn Thr Ala Met Thr Arg Leu Ser
 20 25 30
 Thr Gly Leu Arg Ile Asn Ala Ala Lys Asp Asp Ala Ala Gly Met Ala
 35 40 45
 Ile Gly Glu Lys Met Thr Ala Gln Val Met Gly Leu Asn Gln Ala Ile
 50 55 60
 Arg Asn Ala Gln Asp Gly Lys Asn Leu Val Asp Thr Thr Glu Gly Ala
 65 70 75 80
 His Val Glu Val Ser Ser Met Leu Gln Arg Leu Arg Glu Leu Ala Val
 85 90 95
 Gln Ser Ser Asn Asp Thr Asn Thr Ala Ala Asp Arg Gly Ser Leu Ala
 100 105 110
 Ala Glu Gly Lys Gln Leu Ile Ala Glu Ile Asn Arg Val Ala Glu Ser
 115 120 125
 Thr Thr Phe Asn Gly Met Lys Val Leu Asp Gly Ser Phe Thr Gly Lys
 130 135 140
 Gln Leu Gln Ile Gly Ala Asp Ser Gly Gln Thr Met Ala Ile Asn Val
 145 150 155 160
 Asp Ser Ala Ala Ala Thr Asp Ile Gly Ala His Lys Ile Ser Ser Ala
 165 170 175
 Ser Thr Val Val Ala Asp Ala Ala Leu Thr Asp Thr Thr
 180 185

<210> SEQ_ID NO 70
 <211> LENGTH: 175
 <212> TYPE: PRT
 <213> ORGANISM: Xenorhabdus nematophila

<400> SEQUENCE: 70

Met Ala Ser Val Ile Asn Thr Asn Asp Ser Ala Leu Leu Ala Gln Asn
 1 5 10 15
 Asn Leu Thr Lys Ser Lys Gly Ile Leu Gly Ser Ala Ile Glu Arg Leu
 20 25 30
 Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
 35 40 45
 Ala Ile Ala Asn Arg Phe Thr Ala Asn Val Lys Gly Leu Thr Gln Ala
 50 55 60
 Ala Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly
 65 70 75 80
 Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Ile Arg Glu Leu Thr
 85 90 95
 Val Gln Ser Glu Asn Gly Ser Asn Ser Lys Ser Asp Leu Asp Ser Ile
 100 105 110
 Gln Lys Glu Val Thr Gln Arg Leu Glu Glu Ile Asp Arg Ile Ser Thr
 115 120 125
 Gln Thr Gln Phe Asn Gly Ile Lys Val Leu Asn Gly Asp Val Thr Glu
 130 135 140
 Met Lys Ile Gln Val Gly Ala Asn Asp Asn Glu Thr Ile Gly Ile Lys
 145 150 155 160
 Leu Gly Lys Ile Asn Ser Glu Lys Leu Asn Leu Lys Glu Phe Ser
 165 170 175

<210> SEQ_ID NO 71
 <211> LENGTH: 175
 <212> TYPE: PRT

-continued

<213> ORGANISM: Proteus mirabilis

<400> SEQUENCE: 71

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Met Ala Gln Val Ile Asn Thr Asn Tyr Leu Ser Leu Val Thr Gln Asn
1           5          10          15

Asn Leu Asn Arg Ser Gln Ser Ala Leu Gly Asn Ala Ile Glu Arg Leu
20          25          30

Ser Ser Gly Met Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
35          40          45

Ala Ile Ala Asn Arg Phe Thr Ser Asn Ile Asn Gly Leu Thr Gln Ala
50          55          60

Ser Arg Asn Ala Asn Asp Gly Ile Ser Val Ser Gln Thr Thr Glu Gly
65          70          75          80

Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Ile Arg Glu Leu Thr
85          90          95

Val Gln Ala Lys Asn Gly Thr Asn Ser Asn Ser Asp Ile Asn Ser Ile
100         105         110

Gln Asn Glu Val Asn Gln Arg Leu Asp Glu Ile Asn Arg Val Ser Glu
115         120         125

Gln Thr Gln Phe Asn Gly Val Lys Val Leu Ser Gly Glu Lys Ser Lys
130         135         140

Met Thr Ile Gln Val Gly Thr Asn Asp Asn Glu Val Ile Glu Phe Asn
145         150         155         160

Leu Asp Lys Ile Asp Asn Asp Thr Leu Gly Val Ala Ser Asp Lys
165         170         175

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<210> SEQ ID NO 72

<211> LENGTH: 200

<212> TYPE: PRT

<213> ORGANISM: Butyrivibrio fibrisolvens

<400> SEQUENCE: 72

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Met Val Val Gln His Asn Met Gln Ala Ala Asn Ala Ser Arg Met Leu
1           5          10          15

Gly Ile Thr Thr Gly Asp Gln Ser Lys Ser Thr Glu Lys Leu Ser Ser
20          25          30

Gly Phe Lys Ile Asn Arg Ala Ala Asp Asp Ala Ala Gly Leu Ser Ile
35          40          45

Ser Glu Lys Met Arg Lys Gln Ile Arg Gly Leu Asp Gln Ala Ser Thr
50          55          60

Asn Ala Ser Asp Gly Ile Ser Ala Val Gln Thr Ala Glu Gly Ala Leu
65          70          75          80

Thr Glu Val His Ser Met Leu Gln Arg Met Asn Glu Leu Ala Val Gln
85          90          95

Ala Ala Asn Gly Thr Asn Ser Glu Ser Asp Arg Ser Ser Ile Gln Asp
100         105         110

Glu Ile Asn Gln Leu Thr Thr Glu Ile Asp Arg Val Ala Glu Thr Thr
115         120         125

Lys Phe Asn Glu Thr Tyr Leu Leu Lys Gly Gly Asn Gly Asp Arg Thr
130         135         140

Val Arg Val Tyr Ala His Asp Ala Gly Leu Val Gly Ser Leu Ser Gln
145         150         155         160

Asn Thr Thr Lys Ala Thr Phe Gln Met Arg Lys Leu Glu Ile Gly Asp
165         170         175

Ser Tyr Thr Ile Gly Gly Thr Thr Tyr Lys Ile Gly Ala Glu Thr Val

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180	185	190
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Lys Glu Ala Met Thr Ala Leu Lys
195 200

<210> SEQ ID NO 73
<211> LENGTH: 177
<212> TYPE: PRT
<213> ORGANISM: Bordetella pertussis

<400> SEQUENCE: 73

Met Ala Ala Val Ile Asn Thr Asn Tyr Leu Ser Leu Val Ala Gln Asn
1 5 10 15

Asn Leu Asn Lys Ser Gln Ser Ala Leu Gly Ser Ala Ile Glu Arg Leu
20 25 30

Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Gln
35 40 45

Ala Ile Ala Asn Arg Phe Thr Ala Asn Val Lys Gly Leu Thr Gln Ala
50 55 60

Ala Arg Asn Ala Asn Asp Gly Ile Ser Ile Ala Gln Thr Thr Glu Gly
65 70 75 80

Ala Leu Asn Glu Ile Asn Asn Leu Gln Arg Ile Arg Glu Leu Thr
85 90 95

Val Gln Ala Ser Asn Gly Thr Asn Ser Ala Ser Asp Ile Asp Ser Ile
100 105 110

Gln Gln Glu Val Asn Gln Arg Leu Glu Glu Ile Asn Arg Ile Ala Glu
115 120 125

Gln Thr Asp Phe Asn Gly Ile Lys Val Leu Lys Ser Asn Ala Thr Asp
130 135 140

Met Thr Leu Ser Ile Gln Val Gly Ala Lys Asp Asn Glu Thr Ile Asp
145 150 155 160

Ile Lys Ile Asp Arg Asn Ser Asn Trp Asn Leu Tyr Asp Ala Val Gly
165 170 175

Thr

<210> SEQ ID NO 74
<211> LENGTH: 167
<212> TYPE: PRT
<213> ORGANISM: Clostridium chauvoei

<400> SEQUENCE: 74

Met Ile Ile Asn His Asn Met Asn Ala Leu Asn Ala His Arg Asn Met
1 5 10 15

Met Gly Asn Ile Ala Thr Ala Gly Lys Ser Met Glu Lys Leu Ser Ser
20 25 30

Gly Leu Arg Ile Asn Arg Ala Gly Asp Asp Ala Ala Gly Leu Ala Ile
35 40 45

Ser Glu Lys Met Arg Gly Gln Ile Arg Gly Leu Asp Gln Ala Ser Arg
50 55 60

Asn Ala Gln Asp Gly Ile Ser Leu Ile Gln Thr Ala Glu Gly Ala Leu
65 70 75 80

Ala Glu Thr His Ser Ile Leu Gln Arg Met Arg Glu Leu Ser Val Gln
85 90 95

Ser Ala Asn Asp Thr Asn Val Ala Val Asp Arg Thr Ala Ile Gln Asp
100 105 110

Glu Ile Asn Ser Leu Thr Glu Glu Ile Asn Arg Ile Ser Gly Asp Thr
115 120 125

-continued

Glu Phe Asn Thr Gln Lys Leu Leu Asp Gly Gly Phe Lys Gly Glu Phe
 130 135 140
 Gln Ile Gly Ala Asn Ser Asn Gln Thr Val Lys Leu Asp Ile Gly Asn
 145 150 155 160
 Met Ser Ala Ala Ser Leu Gly
 165

<210> SEQ ID NO 75
 <211> LENGTH: 178
 <212> TYPE: PRT
 <213> ORGANISM: Xanthomonas campestris
 <400> SEQUENCE: 75

Met Ala Gln Val Ile Asn Thr Asn Val Met Ser Leu Asn Ala Gln Arg
 1 5 10 15
 Asn Leu Asn Thr Asn Ser Ser Met Ala Leu Ser Ile Gln Gln Leu
 20 25 30
 Ser Ser Gly Lys Arg Ile Thr Ser Ala Ser Val Asp Ala Ala Gly Leu
 35 40 45
 Ala Ile Ser Glu Arg Phe Thr Thr Gln Ile Arg Gly Leu Asp Val Ala
 50 55 60
 Ser Arg Asn Ala Asn Asp Gly Ile Ser Leu Ala Gln Thr Ala Glu Gly
 65 70 75 80
 Ala Met Val Glu Ile Gly Asn Asn Leu Gln Arg Ile Arg Glu Leu Ser
 85 90 95
 Val Gln Ser Ala Asn Ala Thr Asn Ser Ala Thr Asp Arg Glu Ala Leu
 100 105 110
 Asn Ser Glu Val Lys Gln Leu Thr Ser Glu Ile Asp Arg Val Ala Asn
 115 120 125
 Gln Thr Ser Phe Asn Gly Thr Lys Leu Leu Asn Gly Asp Phe Ser Gly
 130 135 140
 Ala Leu Phe Gln Val Gly Ala Asp Ala Gly Gln Thr Ile Gly Ile Asn
 145 150 155 160
 Ser Ile Val Asp Ala Asn Val Asp Ser Leu Gly Lys Ala Asn Phe Ala
 165 170 175
 Ala Ser

<210> SEQ ID NO 76
 <211> LENGTH: 161
 <212> TYPE: PRT
 <213> ORGANISM: Nitrosomonas europaea
 <400> SEQUENCE: 76

Met Pro Gln Val Ile Asn Thr Asn Ile Ala Ser Leu Asn Ala Gln Arg
 1 5 10 15
 Asn Leu Asn Val Ser Gln Asn Ser Leu Ser Thr Ala Leu Gln Arg Leu
 20 25 30
 Ser Ser Gly Leu Arg Ile Asn Ser Ala Lys Asp Asp Ala Ala Gly Leu
 35 40 45
 Ala Ile Ser Glu Arg Met Thr Ser Gln Ile Arg Gly Met Asn Gln Ala
 50 55 60
 Ala Arg Asn Ala Asn Asp Gly Ile Ser Leu Ala Gln Thr Ala Glu Gly
 65 70 75 80
 Ala Leu Val Glu Ile Gly Asn Asn Leu Gln Arg Ile Arg Glu Leu Ala
 85 90 95

-continued

Val Gln Ser Ala Asn Ala Thr Asn Ser Glu Asp Asp Arg Glu Ala Leu
100 105 110

Gln Lys Glu Val Thr Gln Leu Ile Asp Glu Ile Gln Arg Val Gly Glu
115 120 125

Gln Thr Ser Phe Asn Gly Thr Lys Leu Leu Asp Gly Ser Phe Ala Ser
130 135 140

Gln Ile Phe Gln Val Gly Ala Asn Glu Gly Glu Thr Ile Asp Phe Thr
145 150 155 160

Asp

<210> SEQ ID NO 77

<211> LENGTH: 178

<212> TYPE: PRT

<213> ORGANISM: Campylobacter lari

<400> SEQUENCE: 77

Gly Phe Arg Ile Asn Thr Asn Gly Ala Ser Leu Asn Ala Gln Val Asn
1 5 10 15

Ala Gly Leu Asn Ser Arg Asn Leu Asp Ser Ser Leu Ala Arg Leu Ser
20 25 30

Ser Gly Leu Arg Ile Asn Ser Ala Ala Asp Asp Ala Ser Gly Leu Ala
35 40 45

Ile Ala Asp Ser Leu Lys Thr Gln Ala Asn Ser Leu Gly Gln Ala Ile
50 55 60

Asn Asn Ala Asn Asp Ala Asn Ser Met Leu Gln Ile Ala Asp Lys Ala
65 70 75 80

Met Asp Glu Gln Leu Lys Ile Leu Asp Thr Ile Lys Val Lys Ala Thr
85 90 95

Gln Ala Ala Gln Asp Gly Gln Thr Ala Lys Thr Arg Ala Met Ile Gln
100 105 110

Gly Glu Ile Asn Lys Leu Met Glu Leu Asp Asn Ile Ala Asn Thr
115 120 125

Thr Thr Tyr Asn Gly Lys Gln Leu Leu Ser Gly Ser Phe Ser Asn Ala
130 135 140

Gln Phe Gln Ile Gly Asp Lys Ala Asn Gln Thr Val Asn Ala Thr Ile
145 150 155 160

Gly Ser Thr Asn Ser Ala Lys Val Gly Gln Thr Arg Phe Glu Thr Gly
165 170 175

Ala Val

<210> SEQ ID NO 78

<211> LENGTH: 88

<212> TYPE: PRT

<213> ORGANISM: Salmonella enterica

<400> SEQUENCE: 78

Pro Leu Ala Ser Ile Asp Ser Ala Leu Ser Lys Val Asp Ala Val Arg
1 5 10 15

Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn
20 25 30

Leu Gly Asn Thr Val Thr Asn Leu Asn Ser Ala Arg Ser Arg Ile Glu
35 40 45

Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Lys Ala Gln Ile
50 55 60

Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val Pro
65 70 75 80

-continued

Gln Asn Val Leu Ser Leu Leu Arg
85

<210> SEQ ID NO 79
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: *Pseudomonas aeruginosa*

<400> SEQUENCE: 79

Ala Ile Ala Val Val Asp Asn Ala Leu Ala Ala Ile Asp Ala Gln Arg
1 5 10 15

Ala Asp Leu Gly Ala Val Gln Asn Arg Phe Lys Asn Thr Ile Asp Asn
20 25 30

Leu Thr Asn Ile Ser Glu Asn Ala Thr Asn Ala Arg Ser Arg Ile Lys
35 40 45

Asp Thr Asp Phe Ala Ala Glu Thr Ala Ala Leu Ser Lys Asn Gln Val
50 55 60

Leu Gln Gln Ala Gly Thr Ala Ile Leu Ala Gln Ala Asn Gln Leu Pro
65 70 75 80

Gln Ala Val Leu Ser Leu Leu Arg
85

<210> SEQ ID NO 80
<211> LENGTH: 89
<212> TYPE: PRT
<213> ORGANISM: *Legionella pneumophila*

<400> SEQUENCE: 80

Ala Ile Lys Arg Ile Asp Ala Ala Leu Asn Ser Val Asn Ser Asn Arg
1 5 10 15

Ala Asn Met Gly Ala Leu Gln Asn Arg Phe Glu Ser Thr Ile Ala Asn
20 25 30

Leu Gln Asn Val Ser Asp Asn Leu Ser Ala Ala Arg Ser Arg Ile Gln
35 40 45

Asp Ala Asp Tyr Ala Ala Glu Met Ala Ser Leu Thr Lys Asn Gln Ile
50 55 60

Leu Gln Gln Ala Gly Thr Ala Met Leu Ala Gln Ala Asn Ser Leu Pro
65 70 75 80

Gln Ser Val Leu Ser Leu Leu Gly Arg
85

<210> SEQ ID NO 81
<211> LENGTH: 89
<212> TYPE: PRT
<213> ORGANISM: *Escherichia coli*

<400> SEQUENCE: 81

Pro Leu Glu Thr Ile Asp Lys Ala Leu Ala Lys Val Asp Asn Leu Arg
1 5 10 15

Ser Asp Leu Gly Ala Val Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn
20 25 30

Leu Gly Asn Thr Val Asn Asn Leu Ser Ser Ala Arg Ser Arg Ile Glu
35 40 45

Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Arg Ala Gln Ile
50 55 60

Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Thr Thr
65 70 75 80

-continued

Gln Asn Val Leu Ser Leu Leu Gln Gly
85

<210> SEQ ID NO 82
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: Serratia marcescens
<400> SEQUENCE: 82

Pro	Leu	Ala	Thr	Leu	Asp	Lys	Ala	Leu	Ala	Gln	Val	Asp	Gly	Leu	Arg
1				5				10				15			

Ser Ser Leu Gly Ala Val Gln Asn Arg Phe Asp Ser Val Ile Asn Asn
20 25 30

Leu Asn Ser Thr Val Asn Asn Leu Ser Ala Ser Gln Ser Arg Ile Gln
35 40 45

Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Arg Ala Asn Ile
50 55 60

Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Ser Thr
65 70 75 80

Gln Asn Val Leu Ser Leu Leu Arg
85

<210> SEQ ID NO 83
<211> LENGTH: 89
<212> TYPE: PRT
<213> ORGANISM: Bacillus subtilis
<400> SEQUENCE: 83

Ala	Leu	Thr	Thr	Ile	Lys	Thr	Ala	Ile	Asp	Thr	Val	Ser	Ser	Glu	Arg
1				5				10			15				

Ala Lys Leu Gly Ala Val Gln Asn Arg Leu Glu His Thr Ile Asn Asn
20 25 30

Leu Gly Thr Ser Ser Glu Asn Leu Thr Ser Ala Glu Ser Arg Ile Arg
35 40 45

Asp Val Asp Met Ala Ser Glu Met Met Glu Tyr Thr Lys Asn Asn Ile
50 55 60

Leu Thr Gln Ala Ser Gln Ala Met Leu Ala Gln Ala Asn Gln Gln Pro
65 70 75 80

Gln Gln Val Leu Gln Leu Leu Lys Gly
85

<210> SEQ ID NO 84
<211> LENGTH: 90
<212> TYPE: PRT
<213> ORGANISM: Leptospira interrogans
<400> SEQUENCE: 84

Val	Ile	Gly	Leu	Ala	Asp	Ala	Ala	Leu	Thr	Lys	Ile	Met	Lys	Gln	Arg
1					5			10			15				

Ala Asp Met Gly Ala Tyr Tyr Asn Arg Leu Glu Tyr Thr Ala Lys Gly
20 25 30

Leu Met Gly Ala Tyr Glu Asn Met Gln Ala Ser Glu Ser Arg Ile Arg
35 40 45

Asp Ala Asp Met Ala Glu Glu Val Val Ser Leu Thr Thr Lys Gln Ile
50 55 60

Leu Val Gln Ser Gly Thr Ala Met Leu Ala Gln Ala Asn Met Lys Pro
65 70 75 80

Asn Ser Val Leu Lys Leu Gln Gln Ile

-continued

85

90

<210> SEQ ID NO 85
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: *Shigella sonnei*

<400> SEQUENCE: 85

```
Pro Leu Ser Lys Leu Asp Glu Ala Leu Ala Lys Val Asp Lys Leu Arg
1           5          10          15
Ser Ser Leu Gly Ala Val Gln Asn Arg Phe Asp Ser Ala Ile Thr Asn
20          25          30
Leu Gly Asn Thr Val Asn Asp Leu Ser Ser Ala Arg Ser Arg Ile Glu
35          40          45
Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Arg Ala Gln Ile
50          55          60
Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Thr Thr
65          70          75          80
Gln Asn Val Leu Ser Leu Leu Arg
85
```

<210> SEQ ID NO 86
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: *Edwardsiella tarda*

<400> SEQUENCE: 86

```
Pro Leu Ala Thr Leu Asp Lys Ala Leu Ser Gln Val Asp Asp Leu Arg
1           5          10          15
Ser Gly Leu Gly Ala Val Gln Asn Arg Phe Asp Ser Val Ile Asn Asn
20          25          30
Leu Asn Ser Thr Val Asn Asn Leu Ser Ala Ser Arg Ser Arg Ile Gln
35          40          45
Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Arg Ala Gln Ile
50          55          60
Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Ser Thr
65          70          75          80
Gln Asn Val Leu Ser Leu Leu Arg
85
```

<210> SEQ ID NO 87
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: *Acidovorax avenae*

<400> SEQUENCE: 87

```
Ala Leu Lys Ile Ile Asp Ala Ala Leu Ser Ala Val Asn Gly Gln Arg
1           5          10          15
Ala Ser Phe Gly Ala Leu Gln Ser Arg Phe Glu Thr Thr Val Asn Asn
20          25          30
Leu Gln Ser Thr Ser Glu Asn Met Ser Ala Ser Arg Ser Arg Ile Gln
35          40          45
Asp Ala Asp Phe Ala Ala Glu Thr Ala Asn Leu Ser Arg Ser Gln Ile
50          55          60
Leu Gln Gln Ala Gly Thr Ala Met Val Ala Gln Ala Asn Gln Leu Pro
65          70          75          80
Gln Gly Val Leu Ser Leu Leu Lys
85
```

<210> SEQ_ID NO 88
 <211> LENGTH: 88
 <212> TYPE: PRT
 <213> ORGANISM: Yersinia pestis

 <400> SEQUENCE: 88

 Pro Leu Glu Thr Leu Asp Asp Ala Ile Lys Gln Val Asp Gly Leu Arg
 1 5 10 15

 Ser Ser Leu Gly Ala Val Gln Asn Arg Phe Glu Ser Ala Val Thr Asn
 20 25 30

 Leu Asn Asn Thr Val Thr Asn Leu Thr Ser Ala Arg Ser Arg Ile Glu
 35 40 45

 Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Arg Ala Gln Ile
 50 55 60

 Leu Gln Gln Ala Gly Thr Ser Val Leu Ser Gln Ala Asn Gln Val Pro
 65 70 75 80

 Gln Thr Val Leu Ser Leu Leu Asn
 85

<210> SEQ_ID NO 89
 <211> LENGTH: 88
 <212> TYPE: PRT
 <213> ORGANISM: Photorhabdus luminescens

 <400> SEQUENCE: 89

 Pro Leu Glu Thr Leu Asp Ser Ala Leu Ala Gln Val Asp Ser Leu Arg
 1 5 10 15

 Ser Ser Leu Gly Ala Ile Gln Asn Arg Leu Glu Ser Thr Val Asn Asn
 20 25 30

 Leu Asn Asn Thr Val Asn Asn Leu Ser Ala Ala Arg Ser Arg Ile Glu
 35 40 45

 Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Arg Gly Gln Ile
 50 55 60

 Leu Gln Gln Ala Gly Thr Ala Val Leu Ala Gln Ala Asn Gln Val Pro
 65 70 75 80

 Gln Asn Val Met Ser Leu Leu Arg
 85

<210> SEQ_ID NO 90
 <211> LENGTH: 89
 <212> TYPE: PRT
 <213> ORGANISM: Rhodobacter sphaeroides

 <400> SEQUENCE: 90

 Ala Ile Gly Val Ile Asp Val Ala Leu Ser Lys Ile Ser Gln Ser Arg
 1 5 10 15

 Ser Glu Leu Gly Ala Val Ser Asn Arg Leu Asp Ser Thr Ile Ser Asn
 20 25 30

 Leu Thr Asn Ile Ser Thr Ser Val Gln Ala Ala Lys Ser Gln Val Met
 35 40 45

 Asp Ala Asp Phe Ala Ala Glu Ser Thr Asn Leu Ala Arg Ser Gln Ile
 50 55 60

 Leu Ser Gln Ala Ser Thr Ala Met Leu Ala Gln Ala Asn Ser Ser Lys
 65 70 75 80

 Gln Asn Val Leu Ser Leu Leu Arg Gly
 85

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<210> SEQ ID NO 91
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: Xenorhabdus nematophila

<400> SEQUENCE: 91

```

Pro Leu Asp Thr Leu Asp Lys Ala Leu Ala Gln Val Asp Asp Met Arg
1           5          10          15

Ser Ser Leu Gly Ala Val Gln Asn Arg Leu Glu Ser Thr Val Asn Asn
20          25          30

Leu Asn Asn Thr Val Asn Asn Leu Ser Ala Ala Arg Ser Arg Ile Glu
35          40          45

Asp Ala Asp Tyr Ala Val Glu Val Ser Asn Met Ser Arg Gly Gln Ile
50          55          60

Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val Pro
65          70          75          80

Gln Thr Val Leu Ser Leu Leu Arg
85

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<210> SEQ ID NO 92
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: Proteus mirabilis

<400> SEQUENCE: 92

```

Ala Leu Ala Thr Leu Asp Asn Ala Ile Ser Lys Val Asp Glu Ser Arg
1           5          10          15

Ser Lys Leu Gly Ala Ile Gln Asn Arg Phe Gln Ser Thr Ile Asn Asn
20          25          30

Leu Asn Asn Thr Val Asn Asn Leu Ser Ala Ser Arg Ser Arg Ile Leu
35          40          45

Asp Ala Asp Tyr Ala Thr Glu Val Ser Asn Met Ser Lys Asn Gln Ile
50          55          60

Leu Gln Gln Ala Gly Thr Ala Val Leu Ala Gln Ala Asn Gln Val Pro
65          70          75          80

Gln Thr Val Leu Ser Leu Leu Arg
85

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<210> SEQ ID NO 93
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: Butyrivibrio fibrisolvens

<400> SEQUENCE: 93

```

Ala Ile Asp Ala Ile Ser Asp Ala Leu Ala Lys Val Ser Ala Gln Arg
1           5          10          15

Ser Ala Leu Gly Ser Ile Gln Asn Arg Leu Glu His Ser Ile Ala Asn
20          25          30

Leu Asp Asn Val Val Glu Asn Thr Asn Ala Ala Glu Ser Arg Ile Arg
35          40          45

Asp Thr Asp Met Ala Asp Glu Met Val Thr Tyr Ser Lys Asn Asn Ile
50          55          60

Leu Met Gln Ala Gly Gln Ser Met Leu Ala Gln Ala Asn Gln Ala Thr
65          70          75          80

Gln Gly Val Leu Ser Ile Leu Gln
85

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<210> SEQ ID NO 94
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: Bordetella pertussis

<400> SEQUENCE: 94

```

Ala Leu Ser Lys Leu Asp Asp Ala Met Lys Ala Val Asp Glu Gln Arg
1           5          10          15

Ser Ser Leu Gly Ala Ile Gln Asn Arg Phe Glu Ser Thr Val Ala Asn
20          25          30

Leu Asn Asn Thr Ile Thr Asn Leu Ser Ala Ala Arg Ser Arg Ile Glu
35          40          45

Asp Ser Asp Tyr Ala Thr Glu Val Ser Asn Met Thr Lys Asn Gln Ile
50          55          60

Leu Gln Gln Ala Gly Thr Ser Val Leu Ala Gln Ala Asn Gln Val Pro
65          70          75          80

Gln Asn Val Leu Ser Leu Leu Arg
85

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<210> SEQ ID NO 95
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: Clostridium chauvoei

<400> SEQUENCE: 95

```

Ser Ile Lys Thr Ile Asn Ser Ala Ile Glu Gln Val Ser Thr Gln Arg
1           5          10          15

Ser Lys Leu Gly Ala Val Gln Asn Arg Leu Glu His Thr Ile Asn Asn
20          25          30

Leu Asn Thr Ser Ser Glu Asn Leu Thr Ala Ala Glu Ser Arg Val Arg
35          40          45

Asp Val Asp Met Ala Lys Glu Met Met Ala Phe Ser Lys Asn Asn Ile
50          55          60

Leu Ser Gln Ala Ala Gln Ala Met Leu Gly Gln Ala Asn Gln Gln Pro
65          70          75          80

Gln Gly Val Leu Gln Leu Leu Arg
85

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<210> SEQ ID NO 96
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: Xanthomonas campestris

<400> SEQUENCE: 96

```

Ala Leu Glu Ile Val Asp Lys Ala Leu Thr Ser Val Asn Ser Ser Arg
1           5          10          15

Ala Asp Met Gly Ala Val Gln Asn Arg Phe Thr Ser Thr Ile Ala Asn
20          25          30

Leu Ala Ala Thr Ser Glu Asn Leu Thr Ala Ser Arg Ser Arg Ile Ala
35          40          45

Asp Thr Asp Tyr Ala Lys Thr Thr Ala Glu Leu Thr Arg Thr Gln Ile
50          55          60

Leu Gln Gln Ala Gly Thr Ala Met Leu Ala Gln Ala Lys Ser Val Pro
65          70          75          80

Gln Asn Val Leu Ser Leu Leu Gln
85

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<210> SEQ ID NO 97

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<211> LENGTH: 84
<212> TYPE: PRT
<213> ORGANISM: Nitrosomonas europaea

<400> SEQUENCE: 97

Ile Asp Asp Ala Leu Lys Ile Val Asn Ser Thr Arg Ala Asp Leu Gly
1           5          10          15

Ala Ile Gln Asn Arg Phe Ser Ser Ala Ile Ala Asn Leu Gln Thr Ser
20          25          30

Ala Glu Asn Leu Ser Ala Ser Arg Ser Arg Ile Gln Asp Ala Asp Phe
35          40          45

Ala Ala Glu Thr Ala Ala Leu Thr Arg Ala Gln Ile Leu Gln Gln Ala
50          55          60

Gly Val Ala Met Leu Ser Gln Ala Asn Ala Leu Pro Asn Asn Val Leu
65          70          75          80

Ser Leu Leu Arg

```

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<210> SEQ ID NO 98
<211> LENGTH: 88
<212> TYPE: PRT
<213> ORGANISM: Campylobacter lari

<400> SEQUENCE: 98

Val Met Asp Ile Ala Asp Thr Ala Ile Ala Asn Leu Asp Thr Ile Arg
1           5          10          15

Ala Asn Ile Gly Ala Thr Gln Asn Gln Ile Thr Ser Thr Ile Asn Asn
20          25          30

Ile Ser Val Thr Gln Val Asn Val Lys Ala Ala Glu Ser Gln Ile Arg
35          40          45

Asp Val Asp Phe Ala Ser Glu Ser Ala Asn Tyr Ser Lys Ala Asn Ile
50          55          60

Leu Ala Gln Ser Gly Ser Tyr Ala Met Ala Gln Ala Asn Ala Ala Ser
65          70          75          80

Gln Asn Val Leu Arg Leu Leu Gln
85

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The invention claimed is:

1. A composition comprising a *Salmonella* flagellin 45 polypeptide, wherein the polypeptide comprises an amino acid sequence selected from SEQ ID NOS: 12, 30, 32, 34, 36, 38, 40, 42, and 44.
2. The composition of claim 1, wherein the composition further comprises a radioprotectant.
3. The composition of claim 2, wherein the radioprotectant is selected from an antioxidant, amifostine, vitamin E, a cytokine, a stem cell factor, a growth factor, keratinocyte growth factor, a steroid, 5-androstanediol, and ammonium trichloro(dioxoethylene-O,O')tellurate.
4. The composition of claim 1, wherein the polypeptide is capable of inducing NF κ B activity.
5. A pharmaceutical composition comprising a *Salmonella* flagellin polypeptide, wherein the polypeptide comprises an amino acid sequence selected from SEQ ID NOS: 12, 30, 32, 34, 36, 38, 40, 42, and 44 and a pharmaceutically acceptable adjuvant diluent, or carrier.
6. The pharmaceutical composition of claim 5, wherein the composition further comprises a radioprotectant.
7. The pharmaceutical composition of claim 6, wherein the radioprotectant is selected from an antioxidant, amifostine, vitamin E, a cytokine, a stem cell factor, a growth factor, keratinocyte growth factor, a steroid, 5-androstanediol, and ammonium trichloro(dioxoethylene-O,O')tellurate.
8. The pharmaceutical composition of claim 5, wherein the polypeptide is capable of inducing NF κ B activity.

* * * * *